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(54) Novel pharmacologically active compounds

(57) Novel compounds of the formula:

wherein X is Sor SO and R1, R2, R3, R4, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup>, R<sup>8</sup> and R<sup>15</sup> are organic residues, pharmaceutical compositions containing such compounds particularly for use in the treatment of gastric disorders.

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#### **SPECIFICATION**

## Novel pharmacologically active compounds

The object of the present invention is to provide novel compounds, and therapeutically acceptable salts thereof, which inhibit exogenously or endogenously stimulated gastric acid secretion and provide gastrointestinal cytoprotective effects and thus can 10 be used in the prevention and treatment of peptic

ulcer.

The present invention relates to the use of the compounds of the invention or therapeutically acceptable salts thereof, for inhibiting gastric acid secretion as well as providing gastrointestinal cytoprotective effects in mammals and man. In a more general sense, the compounds of the invention may be used for prevention and treatment of gastrointestinal inflammatory diseases in mammals and man,

- 20 including e.g. gastritis, gastric ulcer, and duodenal ulcer. Furthermore, the compounds may be used for prevention and treatment of other gastrointestinal disorders, where cytoprotective and/or gastric antisecretory effect is desirable e.g. in patients with
- 25 gastrinomas, in patients with acute upper gastrointestinal bleeding, and in patients with a history of chronic and excessive ethanol consumption. The invention also relates to pharmaceutical compositions containing at least one compound of the

30 invention, or a therapeutically acceptable salt thereof, as active ingredient. In a further aspect, the invention relates to processes for preparation of such new compounds and to novel intermediates in the preparation of the compounds of the invention.

Benzimidazole derivatives intended for inhibiting gastric acid secretion are disclosed in the British patent specifications 1 500 043 and 1 525 958, in the US patent 4 182 766, in the European patent specification 0 005 129, and in the Belgian patent specifica-

- 40 tion 890 024. Benzimidazole derivatives proposed for use in the treatment or prevention of special gastrointestinal inflammatory disease are disclosed in the European patent application with publication no. 0 045 200.
- It has been found that the compounds of the formula

wherein

R15 is H, CH3 or C2H5;

- 50 R1, R2, R3 and R4, which are the same or different, are
  - (a) H
  - (b) halogen
  - (c) —CN
  - (d) -CHO
- 55 (e) -CF<sub>3</sub>

(h) --CH(OR13)2

- (i)  $-(Z)_n A D$
- 60 (j) aryl
  - (k) aryloxy
    - (I) alkylthio containing 1-6 carbon atoms
  - (m) --NO<sub>2</sub>
  - (n) alkylsulfinyl containing 1-6 carbon atoms
- 65 or wherein
  - (o) adjacent groups R1, R2, R3 and R4 together with the adjacent carbon atoms in the benzimidazole ring form a 5-, 6- or 7-membered monocyclic ring or a 9-, 10- or 11-membered bicyclic ring which rings may be saturated or unsaturated and may contain 0-3 hetero atoms selected from N and O, and which rings may be optionally substituted with 1-4 substituents selected
- from alkyl groups with 1-3 carbon atoms, alkylene radicals containing 4-5 carbon atoms giving spiro
- 75 compounds, or two or four of these substituents together form one or two oxo groups 0
- (-C-), whereby if R1, R2, R3 and R4 together with the adjacent carbon atoms in the benzimidazole ring form two rings they may be condensed with each other, in which formulas R11 and R12, which are the same or different, are
  - (a) aryl,
  - (b) alkoxy containing 1-4 carbon atoms,
  - (c) alkoxyalkoxy containing 1-3 carbon atoms in each alkoxy part.
- (d) arylalkoxy containing 1-2 carbon atoms in the
  - alkoxy part,
    - (e) aryloxy,
- (f) dialkylamino containing 1-3 carbon atoms in the
- (g) pyrrolidino or piperidino, optionally substituted with alkyl containing 1-3 carbon atoms
  - R<sup>13</sup> is (a) alkyl containing 1-4 carbon atoms, or (b) alkylene containing 2-3 carbon atoms:

nis0or1;

- A is (a) alkylene containing 1-6 carbon atoms
- (b) cycloalkylene containing 3-6 carbon atoms
- (c) alkenylene containing 2-6 carbon atoms
- 100 (d) cycloalkenylene containing 3-6 carbon atoms,
  - (e) alkynylene containing 2-6 carbon atoms; Dis(a) -

(c)  $-(Y)_m - (C)_r - R^{10}$ 

wherein

R9 is (a) alkoxy containing 1-5 carbon atoms, or

(b) dialkylamino containing 1-3 carbon atoms in the alkyl parts;

mis0or1;

ris0or1;

Yis (a) -0-

(b) -NH-(c) -NR10-:

R10 is (a) H

(b) alkyl containing 1-3 carbon atoms,

(c) arylalkyl containing 1-2 carbon atoms in the alkyl part, or

(d) aryl:

R5 is (a) Hor

(b) ---C-

15 wherein

R14 is (a) alkyl containing 1-6 carbon atoms,

(b) arylalkyl containing 1-2 carbon atoms in the alkyl part

(c) aryl

(d) alkoxy containing 1-4 carbon atoms

(e) arylalkoxy containing 1-2 carbon atoms in the alkyl part

(f) aryloxy

(g) amino

(h) mono- or dialkylamino containing 1-4 carbon atoms in the alkyl part(s)

(i) arylalkylamino containing 1-2 carbon atoms in the alkyl part

(j) arylamino;

R<sup>6</sup> and R<sup>8</sup>, which are the same or different, are

(a) Hor

(b) alkyl containing 1-5 carbon atoms;

R7is(a) H

(b) alkyl containing 1-8 carbon atoms

(c) alkoxy containing 1-8 carbon atoms

(d) alkenyloxy containing 2-5 carbon atoms

(e) alkynyloxy containing 2-5 carbon atoms

(f) alkoxyalkoxy containing 1-2 carbon atoms in each alkoxy group

(g) dialkylaminoalkoxy containing 1-2 carbon atoms in the alkyl substituents on the amino nitrogen and 1-4 carbon atoms in the alkoxy group

(h) oxacycloalkyl containing one oxygen atom and 3-7 carbon atoms

(i) oxacycloalkoxy containing two oxygen atoms and 4-7 carbon atoms

(j) oxacycloalkylalkyl containing one oxygen atom

and 4-7 carbon atoms (k) oxacycloalkylalkoxy containing two oxygen

50 atoms and 4-6 carbon atoms, or

(I) R<sup>6</sup> and R<sup>7</sup>, or R<sup>7</sup> and R<sup>8</sup> together with the adjacent carbon atoms in the pyridine ring form a ring wherein the part constituted by  $R^6$  and  $R^7$ , or  $R^7$  and R<sup>8</sup>, is

-CH=CH-CH=CH 55

-O---(CH₂)<sub>p</sub>---

---CH<sub>2</sub>(CH<sub>2</sub>)<sub>p</sub>---

-O-CH=CH-

-NH-CH=CH-

-N--CH=CH---

wherein p is 2, 3 or 4 and the 0 and N atoms always

are attached to position 4 in the pyridine ring; and physiologically acceptable salts of the com-

pounds I wherein X is S;

with the provisos that

(a) not more than one of R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> is hydrogen,

(b) when X is SO,  $R^5$  is H and  $R^6$ ,  $R^7$  and  $R^8$  are selected only from hydrogen, methyl, methoxy,

70 ethoxy, methoxyethoxy and ethoxyethoxy and at the same time more than one of R1, R2, R3 and R4 are hydrogen, then R1, R2, R3 and R4 cannot be selected only from alkyl groups, halogen, alkoxycarbonyl, alkoxy or alkanoyl,

(c) when X is S, R5 is H, alkanoyl or alkoxycarbonyl, and R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> are selected only from hydrogen, methyl, ethyl, methoxy, ethoxy, methoxyethoxy and ethoxyethoxy and at the same time more than one of  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  are hydrogen, then  $R^7$ ,  $R^2$ ,  $R^3$  and  $R^4$ cannot be selected only from alkyl groups, halogen, alkoxycarbonyl, alkoxy, alkanoyl, trifluormethyl, or

NO<sub>2</sub>

(d) when X is SO, one of R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> is H and the other two of R6, R7 and R8 are alkyl, and at the same time more than one of R1, R2, R3 and R4 are hydrogen, then those radicals R1, R2, R3 and R4 which are not H cannot be selected only from alkyl, halogen, cyano,

-(alkoxy), (alkyl) — OC — (alkyl —, alkoxy,

hydroxyalkyl, CF3, or (alkyl) — C —,

(e) when R3, R4, R5 and R15 are H and simultaneously R<sup>6</sup> and R<sup>8</sup> are H or CH<sub>3</sub> and R<sup>7</sup> is OCH<sub>3</sub>, then R<sup>1</sup> is not CF3 when R2 is H, and R2 is not CF3 when R1 is H, are effective as gastrointestinal cytoprotectives and as inhibitors of gastric acid secretion in mammals and man as stated above.

Illustrative examples of the various radicals in the formula lare as follows. These illustrative examples will be applicable to different radicals depending on the number of carbon atoms prescribed for each

100 radical. It will be understood that the expressions "alky!" and "alkoxy" include straight, branched and cyclic structures.

· TVD.

F. Cl. Br. I

Alwyl: CH3, C2H5, n-C3H7, 1-C3H7, n-C4Hg, sec.-C4Hg,

iso.- $C_4H_9$ , tert.- $C_4H_9$ , n- $C_5H_{11}$ , n- $C_6H_{13}$ ;

-CH<sub>2</sub>+, -CH<sub>2</sub>CH<sub>2</sub>+, -(CH<sub>2</sub>)<sub>3</sub>+, -CH<sub>2</sub>-CH+ , -(Ch<sub>2</sub>)<sub>4</sub>+,

Alkenylenes -CH+CH- . -CH2+CH+CH-. . -CH2-CH+CH-CH2+ . -(cH<sub>2</sub>)<sub>2</sub>-CH-CH-CH<sub>2</sub>- , -(CH<sub>2</sub>)<sub>3</sub>-CH-CH-CH<sub>2</sub>-

-S-CH3, -S-C2H5, -S-1-C3H7 Alkylthio:

-с-с- , -сн<sub>2</sub>-с-с- . Alkynylene:

-OCH3 . -OC2H5 . -O-n-C3H7 . -O-1-C3H7 .

-O-n-C<sub>4</sub>M<sub>g</sub> , -O-isa-C<sub>4</sub>H<sub>g</sub> , -O-sec.-C<sub>4</sub>H<sub>g</sub> ,

-O-tert.-C4Hg, -O-n-C5H11 .

-осн<sub>2</sub>осн<sub>3</sub> , -осн<sub>2</sub>сн<sub>2</sub>осн<sub>3</sub>, -осн<sub>2</sub>сн<sub>2</sub>осн<sub>2</sub>сн<sub>3</sub>,

-0CH2CH2CH2OCH2CH2CH3

Aryloxyı

-0-CH-CH<sub>2</sub> , -0-CH-CH-CH<sub>3</sub> , -0-CH-CH-С<sub>2</sub>H<sub>5</sub>. -0-CH2-CH-CH-CH2CH3

Alkynylaky: -0-CMCH. -0-CMg-CMCH. -0-CMg-CMC+CMg -0-CMg-C=2-CMgCMg

i...strative examples of the radical  $\neg \mathsf{CH}(\mathsf{OR}^{13})_{\mathbb{Z}}$  are:

Illustrative examples of the ring structures involving  $\sigma^{2}_{\ \mu}$   $R^{2}$  ,  $R^{3}$  or  $R^{4}$  are

. muere v is

-CH2CH2CH2-CH2CH2CH2CH2-CH2-C(CH3)2-CH2-(CH2)5-CH-CH-CH-CH-CH3
-CH-CH2CH2-CH3-CH-CH2-CH3-CH-CH2-CH3-CH3-CH3-CH3-CH3-CH3-CH3-CH3-CH3-CH2)2-NH-OCH2O-OCH2CH2O-OC(CH2)30-

CH<sub>2</sub> CH<sub>3</sub>

The radical -(Z) $_{\rm n}$ - A - O comprises the following radicals. The expression (alkyl 1-3g) etc. means alkyl groups containing 1, 2 or 3 carbon atoms.

A - CN

A - C - O -(alkyl 1-5c)

A - C - N (alkyl 1-3c)

(alkyl 1-3c)

A - H

A - (alkyl 1-3c)

A - (alkyl 1-2c)-aryl

A - aryl

A - 0 - H

A - 0 - (alkyl 1-3c)

A - 0 - (alkyl 1-3c)

A - 0 - (alkyl 1-3c)

A - 0 - aryl

A - NH - H

A - NH - [alkyl 1-3c]

A - HH - aryl

gl0

A - N - H

gl0

A - N - H R10 A - N - (elbyl l-3cl R10 A - N - (elbyl l-2c)-eryl R10 A - N - eryl A - O - C - H

A- 0 - C - (alkyl 1-3c)
A- 0 - C - (alkyl 1-2c)-aryl
O
A- 3 - C - aryl

A- NH - C -(alkyl 1-3c) Q A- NH - C -(alkyl 1-2c)-aryl

A- NH - eryl
R10 0
A- N - C - H
R10 0
A - N - C - (elkyl 1-3c)
R10 0
A - N - C - (elkyl 1-2c)-eryl
R10 0
A - N - C - eryl
-0 -A - CN
-0 -A - CN
(elkyl 1-3c)
-0 -A - C - (elkyl 1-3c)
(elkyl 1-3c)
(elkyl 1-3c)

-0-A - H -0-A -(alkyl 1-3c) -0-A-(alkyl 1-2c)-aryl -0-A-aryl

-0 - A - 0 - H -0 - A - 0 - (alkyl 1-3c) -0 - A - 0 - (alkyl 1-2c)-aryl -0 - A - 0 - aryl -0 - A - NH - H -0 - A - NH -(alkyl 1-3c)
-0 - A - NH -(alkyl 1-2c)-aryl
-0 - A - NH - aryl

R10

-0 - A - N - H

R10

-0 - A - N - (alkyl 1-3c)

R10

-0 - A - N -(alkyl 1-2c)-aryl

R10

-0 - A - N - (alkyl 1-2c)-aryl

R10

-0 - A - N - aryl

-0 - A - 0 - C - M

-0 - A - 0 - C - (alkyl 1-3c)

-0 - A - 0 - C - (alkyl 1-2c)-aryl

-0 - A - 0 - C - aryl

-0 - A - NH - C - (alkyl 1-3c)

-0 - A - NH - C - (alkyl 1-3c)

-0 - A - NH - aryl

R10 0

-0 - A - N - C - (alkyl 1-3c)

-0 - A - N - C - (alkyl 1-3c)

-0 - A - N - C - (alkyl 1-3c)

-0 - A - N - C - (alkyl 1-3c)

0 -C- A -CN 0 -C- A -C - O-(alkyl 1-5c) 0 0 0 (alkyl 1-3c) -C- A -C - N (alkyl 1-3c)

-C -A -H

-C -A -(alkyl 1-3c)

-C -A -(alkyl 1-2c)-aryl

-C -A -aryl

-C -A -O -H

-C -A -O -(alkyl 1-3c)

-C -A -O -(alkyl 1-2c)-aryl

-C -A -O -aryl

-C -A -NH -H

-C -A -NH -(alkyl 1-3c)

-C -A -NH -(alkyl 1-2c) -aryl

-C -A -NH -aryl

-C -A -N -H

-C -A -N -aryl

-C -A -N -(alkyl 1-3c)

0 g
-C-A-NH-C-(alkyl 1-2C)-aryl
3
-C-A-NH-aryl

о R<sup>10</sup> о -C-A-N - C-H

0 R 0 0 H I H -C-A-N - C-(alkyl 1-3C)

o R<sup>10</sup> g -C-A-N - C-aryl

The radical -C-R<sup>11</sup> comprises the following radicals.

-C-aryl

-C-0-(alkyl 1-4C)

-C-0-(alkyl 1-3 c)-0-(alkyl 1-3c)

-C-0-(alkyl 1-2c)-aryl

-C-0-aryl

-C-N (alkyl 1-3c)

(alkyl 1-3c)

-C-N (optionally substituted with alkyl)

-C-N [optionally substituted with alkyl)

The radical -0-c-R<sup>12</sup> comprises the following radicals.

-0-c-aryl

0
-0-c-0-(alkyl 1-4c)

-0-c-0-(alkyl 1-3c)-0-(alkyl 1-3c)

-0-c-0-aryl

0
-0-c-0-aryl

(alkyl 1-3c)

-0-c-n

(alkyl 1-3c)

(optionally substituted with alkyl)

-O-C-N (optionally substituted with alkyl)
-O-C-N (optionally substituted with alkyl)

The radical -C-R<sup>14</sup> comprises the following radicals:

O
-C-(alkyl 1-6c)

-c-(alkyl 1-6c)
0
-c-(alkyl 1-2c)-aryl
0
-c-aryl
0
-c-0-(alkyl 1-4c)
0
-c-0-(alkyl 1-2c)-aryl

0
-c-O-aryl
0
-c-NH2
0
-c-NH(alkyl 1-4c)
0
-c-N(alkyl 1-4c)
-c-N(alkyl 1-4c)
0
(alkyl 1-4c)
0
(alkyl 1-4c)
0
-c-N
aryl
0
-c-N(aryl)

b.

Further illustrative examples of the radicals in the formula I are:

alkylsulfinyl:

SOCH3. SOC2H5. SOCH2CH2CH3. SO-4-C3H7. SO-n-C4H6. SO-n-C5H11

oxacycloalkyl:

$$\bigcirc$$

oxacycloalkoxy:

oxecycloalkyl-alkyl:

oxacycloalkyl-alkoxy:

The compounds of the invention that are sulfoxides (X=SO) have an asymmetric centre in the sulfur atom, i.e. these compounds exist as two optical isomers (enantiomers), or if they also contain one or 5 more asymmetric carbon atoms the compounds have two or more diastereomeric forms, each existing in two enantiomeric forms. Such asymmetric carbon atoms may be the carbon atom on which R<sup>15</sup> is attached (when R<sup>15</sup> is other than H) or a carbon atom 10 in some of the substituents.

Both the pure enantiomers, racemic mixtures (50% of each enantiomer) and unequal mixture of the two are within the scope of the present invention. It should be understood that all the diastereomeric 15 forms possible (pure enantioners or racemic mix-

tures) are within the scope of the invention.

The compounds of the invention that are sulfides (X=S) may be asymmetric due to one or more asymmetric carbon afoms, as described above. The 20 different diasetereomeric forms possible as well as the pure enantiomers and racemic mixtures are

within the scope of the invention.
It should be noted that for all the compounds of the invention wherein R<sup>5</sup> is H the substituents R<sup>1</sup> and R<sup>4</sup>
25 as well as R<sup>2</sup> and R<sup>3</sup> are considered to be equivalent.

This is due to the tautomerism in the imidazole part of the benzimidazole nucleus causing an equilibrium between the two possible NH-forms. This is illustrated by the following example:

- 30 I Preferred groups of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are:
  - 1. F
  - halogens F, CI, Br and the groups CN, CHO, CO(aryI), COO(alkyI), CF<sub>3</sub>, SCH<sub>3</sub>, SOCH<sub>3</sub> and NO<sub>2</sub>
- 35 3. the groups alkylene-D, O-alkylene-D and CO-alkylene-D wherein D is CN, COO(alkyl), COR<sup>10</sup>, OR<sup>10</sup> and R<sup>10</sup>
  - 4. aryl and aryloxy

- 40 6. —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—,—CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—and • —CH=CH—CH=CH—
  - 7. —CH=CH-CH=C-(CH<sub>2</sub>)<sub>2-3</sub>--
  - 8. saturated haterocyclic ring structures having 2

45 oxygen atoms.

- 9. unsaturated 6-membered heterocyclic ring structures having one nitrogen atom
- II Further preferred groups of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are:
- 50 1. H
  - 2. halogers Cland Brand the groups CO(phenyl), COOCH<sub>3</sub> CF<sub>3</sub> SCH<sub>3</sub> and SOCH<sub>3</sub>
  - 3. the groups alkyl, alkoxyalkyl, aryloxyalkyl, arylaktyl, aryl
- 4. the groupsalkoxy, alkoxyalkoxy, aryloxyalkoxy, arylalkoxy, aryloxy
  - 5. the group alkanoyl
  - 6. —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—,—CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—and —CH=CH—CH=CH—
- 50 | 7. —CH=<del>CH</del>-CH=C—(CH<sub>2</sub>)<sub>2-3</sub>--
  - 8. saturated beterocyclic ring structures having 2 oxygen atoms in 4,5-,5,6- or 6,7-"catechol positions", e.g. (5,6-position shown)

- 65 III Still further preferred groups of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are:
  - 1. H
  - 2. Brandthegroups COOCH3 and CF3
  - 3. the groups CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub>, CH(CH<sub>3</sub>)<sub>2</sub>, CH<sub>3</sub>OCH<sub>2</sub>CH<sub>2</sub>--,
- 70 phenyl
- 4. the groups CH<sub>3</sub>O, CH<sub>3</sub>(CH<sub>2</sub>)<sub>6</sub>O—, CH<sub>3</sub>OCH<sub>2</sub>CH<sub>2</sub>O—, (phenyl)-OCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>O—, (phenyl)CH<sub>2</sub>CH<sub>2</sub>O—, (phenyl)O—
  - 5. the groups CH<sub>3</sub>CO--, C<sub>2</sub>H<sub>5</sub>CO--
- 75 6. —CH2CH2CH2—, —CH2CH2CH2CH2—
  - 7. —OCH-@-, -0 in the 5,6-"catechol position"
  - IV Particularly preferred groups of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are:
  - H, COOCH<sub>3</sub>, CF<sub>3</sub>, CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub>, CH(CH<sub>3</sub>)<sub>3</sub>, CH<sub>3</sub>O,
- 80 —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>—, —CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>— and —OCH<sub>2</sub>O— V In a preferred embodiment, at least three of the radicals R<sup>1</sup>, B<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are other than hydrogen, or

they form at least one ring.

VI In another preferred embodiment the radicals R1 and R2 form a ring structure

VII In another preferred embodiment the radicals R<sup>2</sup> 5 and R<sup>3</sup> form a ring structure.

VIII In a preferred embodiment at least three of the radicals R1, R2, R3 and R4 are other than hydrogen. IX In a preferred embodiment the radicals R1, R2, R3 and R4 are selected from H, halogen, CF3, alkyl and 10 alkoxy groups.

X In a preferred embodiment the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R4 are selected from H, alkyl and alkoxy groups. XI In a preferred embodiment the radicals R1, R2, R3 and R4 are selected from H and alkyl groups.

15 XII The preferred groups of X is S. XIII The preferred group of X is SO.

VIX The preferred group of R15 is H.

XV Preferred groups of the radical R<sup>5</sup> are H. arylcarbonyl, alkoxycarbonyl, arylalkoxycarbonyl, di-

20 alkylaminocarbonyl and arylaminocarbonyl. XVI Further preferred groups of the radical R5 are H, phenylcarbonyl, methoxycarbonyl, tert-butoxycarbonyl, benzyloxycarbonyl, dimethylaminocarbonyl

and phenylaminocarbonyl. 25 XVII Particularly preferred of the radical R5 is H. XVII Preferred groups of the radicals R<sup>6</sup> and R<sup>8</sup> are: H, CH<sub>3</sub>, C<sub>2</sub>H25, C<sub>3</sub>H<sub>7</sub> and CH(CH<sub>3</sub>)<sub>2</sub>

2. ring structures connecting position 4 in the pyridine ring.

30 XIX Particularly preferred groups of the radicals R6 and R<sup>5</sup> are H, CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub> and ring structures also connecting position 4 in the pyridine ring XX Preferred groups of the radical R7 are:

1. H, CH3, C2H5 35 2. OCH<sub>3</sub>, OC<sub>2</sub>H<sub>5</sub>, OCH<sub>2</sub>CH<sub>2</sub>CH<sub>3</sub>, O(CH<sub>2</sub>)<sub>3</sub>CH<sub>3</sub>, OCH<sub>2</sub>

3. OCH2CH=CH2, OCH2C=CH

4. OCH<sub>2</sub>CH<sub>2</sub>OCH<sub>3</sub>, OCH<sub>2</sub>

5. OCH2CH2N(CH3)2

6. —CH=CH—CH=CH-bound to positions 3 and 4,

-CH=CH--CH=CH-bound to positions 4 and 5, 40

-CH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>-bound to positions 3 and 4,

-CH2CH2CH2-bound to positions 4 and 5, -CH2CH2CH2CH2-bound to positions 3 and 4,

-- CH2CH2CH2CH2-bound to positions 4 and 5,

-OCH<sub>2</sub>CH<sub>2</sub>-bound to positions 3 and 4.

-OCH2CH2-bound to positions 4 and 5.

-OCH2CH2CH2-bound to positions 3 and 4,

-OCH<sub>2</sub>CH<sub>2</sub>CH<sub>2</sub>-bound to positions 4 and 5,

XXI Further preferred groups of the radical R7 are:

50 1. CH<sub>2</sub>

2. OCH<sub>3</sub>, OC<sub>2</sub>H<sub>5</sub>, OCH<sub>2</sub>CH<sub>2</sub>CH(CH<sub>3</sub>)<sub>2</sub>

3. OCH<sub>2</sub>CH=CH<sub>2</sub>

4. OCH<sub>2</sub>CH<sub>2</sub>OCH<sub>3</sub>, OCH<sub>2</sub>,

5. -- CH2CH2CH2-bound to positions 3 and 4,

55 —CH<sub>2</sub>CH<sub>2</sub>-bound to positions 4 and 5,

-CH2CH2CH2CH2-bound to positions 3 and 4,

-CH2CH2CH2CH2-bound to positions 4 and 5,

-OCH2CH2-bound to positions 3 and 4, -OCH2CH2-

bound to positions 4 and 5, —OCH2CH2CH2-bound to

60 positions 3 and 4, —OCH<sub>2</sub>CH<sub>2</sub>-bound to positions 4 and 5.

XXII - Particularly preferred groups of the radical R7 are CH<sub>3</sub>, OCH<sub>3</sub>, OCH<sub>2</sub>CH<sub>2</sub>CH(CH<sub>3</sub>)<sub>2</sub>,—OCH<sub>2</sub> ,

—OCH₂CH₂CH₂-bound to positions 3 and 4 or to 65 positions 4 and 5.

XXIII Preferred pyridyl substitution patterns are:

XXV Still further preferred pyridyl substitution patterns are:

XXVI Particularly preferred pyridyl substitution patterns are:

5 XXVII. In a preferred embodiment two of the radicals  $R^6$ ,  $R^7$  and  $R^6$  form one ring structure and the third radical of R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> is H.

XXVIII In a preferred embodiment R15 and R5 are H, at least three times of the radicals R1, R2, R3 and R4 are 10 other than H, R<sup>6</sup> and R<sup>8</sup> are H or CH<sub>3</sub> and R<sup>6</sup> is CH<sub>3</sub>,

OCH3 or OCH2CH=CH2.

XXIX In a preferred embodiment R15 and R5 are H, the radicals R1, R2, R3 and R4 form at least one ring structure, R<sup>6</sup> and R<sup>8</sup> are H or CH<sub>3</sub> and R<sup>7</sup> is CH<sup>3</sup>, OCH<sup>3</sup> 15 or OCH2CH=CH2.

XXX Preferred compounds are those of the formula

wherein  $R^2$  is alkyl or alkoxy, preferably  $CH_3$ ,  $C_2H_5$ , CH(CH<sub>3</sub>)<sub>2</sub> and OCH<sub>3</sub>, and X is S or SO.

Further illustrative examples of the radicals in the 20 formula lare given in the examples and lists of specific compounds given elsewhere in this specification.

Illustrative examples of compounds included in the scope of the invention are given in the following

Table 1.

Illustrative examples of compounds included in the scope of the invention.

					•				1	
	R <sup>15</sup>	R <sup>3</sup>	R <sup>2</sup>	R3	84	R <sup>S</sup>		R <sup>6</sup>	R <sup>7</sup>	ę\$
	н	CH <sub>3</sub>	013	CH <sup>3</sup>	CH3	и	- 1	Oly	001201-012	CH <sup>3</sup>
٥	м	CH <sub>3</sub>	CH <sub>3</sub>	CH <sub>3</sub>	CH3	н		CN3	001 <sup>2</sup> CH-CH <sup>2</sup>	. CH3
,	и	CH,	CH <sub>3</sub>	CH3	CN <sub>3</sub>	H .		Ol <sub>3</sub>	осн <sub>3</sub>	CH <sub>3</sub>
٥		CH <sub>3</sub>	CH <sub>3</sub>	CH <sub>3</sub>	CH <sub>3</sub>	н	Ξ,	CII3	осн3	ᅄ
,	H	01	СНЗ	CH <sub>3</sub>	н	н		CH3	001201-012	CH <sub>3</sub>
٥		OH,	CH <sub>1</sub>	CH <sub>3</sub>	н	н		CH <sub>3</sub>	OCHZCH-CHZ	. CH <sub>3</sub>
	H	CH <sub>3</sub>	OI <sub>1</sub>	СН3	н	н		CH <sub>3</sub>	OCH <sub>3</sub>	CH <sub>3</sub>
٥		CH <sub>3</sub>	OI <sub>3</sub>	СН	и.	н		CH <sub>3</sub>	осн <sub>3</sub>	. сн <sub>3</sub>
	н	CH,	CH <sub>3</sub>	H	CH <sub>3</sub>	н		CN3	OCHZCH-CHZ	CH3
٥	н	CH <sub>3</sub>	CH3	н	CH <sub>3</sub>	H		CH <sub>3</sub>	OCH2CH+CH2	CH <sub>3</sub>
	н	οι,	CH	н	CH <sub>3</sub>	н		CH3	осн <sub>а</sub>	СНЗ
0	H	Ot <sub>3</sub> .	CH3	н	CH3	H		CH <sub>3</sub>	осн <sub>3</sub>	CH <sub>3</sub>
5	н	ᅄ	CHa	н	н	н		CH <sub>3</sub>	оси <sub>2</sub> си-си <sub>2</sub>	CH <sub>3</sub>
		,	۵.,					•	• •	1

	-15	<b>8</b> <sup>1</sup>	ę²	۲	n4	<b>8</b> 5	A <sup>b</sup>	R <sup>7</sup>	فع
	a <sup>15</sup>	<u> </u>					OI,	OCH <sup>2</sup> CH-CH <sup>2</sup>	۵,
9	4	O-3	o <sub>3</sub>		-		013	OCH,CH-CH,	CH <sub>3</sub>
	16		or <sup>3</sup>	O13		*	CH <sub>3</sub>	מכוליםו-טול	OI,
)	*	H	· 013	α <sub>3</sub>				OCH CH	Di <sub>3</sub>
		α <sub>j</sub>	14	H	Cr3	N	O'3		o's
	×	Ol <sub>3</sub>	•	16	ᅄ	Ħ	CH <sub>3</sub>	001201-012	
		DY.	*	H	×	*	Ol3	OCH <sup>2</sup> CH-CH <sup>2</sup>	OI <sub>3</sub>
		Ol <sub>3</sub>	×		×	H	OI3	001201-013	CH <sub>3</sub>
		, <del>- ,</del>	OL <sub>3</sub>	я	×	H.	013	001201-013	ОĠ
	•		01,	×	и	M	O13	901 <sup>2</sup> 01-01 <sup>3</sup>	ᅄ
)	H		•	21	M	и	CH <sub>3</sub>	0050+05	O13
	H	×	901 <sub>3</sub>	н	×	Ħ	cاc	001501-015	CH <sup>3</sup>
•	H	M	, acri			н	CH <sub>3</sub>	001 <sup>2</sup> C±01	CH <sub>3</sub>
	Ħ	М.	9013				OI <sub>3</sub>	OCHTERN	CH <sub>2</sub>
0	×	H	0CH <sup>3</sup>		-	H	CH <sub>3</sub>	0(012)301-012	о,
0	×	H	0C/3	ĸ	•	. к	O13	0(OL)3Ol3	OI,
٥	ĸ	H	. <sub>001</sub> 3	<b>n</b> .	М		ο <sub>1</sub> 3	0CH(CH <sub>3</sub> ) <sub>2</sub>	CH <sub>2</sub>
,		H	αcι <sub>3</sub>	н	H	H			OI <sub>3</sub>
٥	*	H	0CH <sub>3</sub>	M	и	ĸ	C <sub>IO</sub> 3	00H(CH3)2	-
		11	0013	н		*	C <sup>IO</sup> 3	oc(cH <sup>3</sup> )3	Oly
	<b>H</b>		-	н	*	H	CH3	oc(cH <sub>3</sub> ) 3	CN
50	н	Ħ	осн						cor

					35	16	x <sup>2</sup>	6
A <sup>15</sup>	<b>a</b> 1	R <sup>2</sup>	r <sub>3</sub>	a^	<u> </u>			
		0CH <sub>3</sub>	. н	н	и.	CH3	•♦ . S. 3	Ol
		•	×	и	10	O13	•	Cr3
) H H	*	∞್ಯ ಜ್ಯು	н	×	M	· OI3	0CH2-	<sub>CH</sub> 3
н		00/ <sub>3</sub>	н	×	· N	013	och₂-△	CH3
<b>*</b>		0CK <sup>3</sup>	ж	н	×	CH3	∞1 <sub>2</sub> -<	O13
<b>M</b>	*	ocn <sup>3</sup>	ж.,	H	H	cH3	0CH <sup>2</sup> -<	CH3
) H		001 <sub>3</sub>	×	Ħ	×	cu <sup>3</sup>	0(CH2)2H(CH3)2	O43
<b>K</b>	H	00h	н	н ,	н	OI3	0(012)2191(013)2619	
H	** *	0CH <sub>3</sub>	н	-ж	Ħ	or <sup>3</sup>	0(CH <sup>2</sup> ) <sup>2</sup> H(CH <sup>2</sup> ) <sup>2</sup>	DI3
) #	*	0CH <sub>3</sub>	×	н	H	O*3	001 <sup>2</sup> 01 <sup>2</sup> 01(01 <sup>3</sup> ) <sup>5</sup>	CH <sub>2</sub>
	# H	00K <sub>3</sub>	H	M	* <b>N</b>	Cr <sup>3</sup>	OCH <sup>5</sup> CH <sup>5</sup> CH(CH <sup>3</sup> ) <sup>5</sup>	013
	# #	001 <sub>3</sub>	н	и	N	, N	OCH3	(Z)
) X	, K	och <sub>3</sub>	н	н	×	H	0(042)3043	ري
		00H <sub>3</sub>	н	ж	н	н	6(01 <sup>5</sup> ) <sup>2</sup> 01 <sup>3</sup>	ري
۸ ه	*	оси <sub>3</sub>	×	H	н	CH3	OCH <sup>2</sup> CH <sup>2</sup> CH <sup>2</sup> CH(CH <sup>3</sup> ) <sup>3</sup>	CH <sub>2</sub>
	 Di	0CH <sub>3</sub>	· c+ <sub>3</sub>	×	×	*	cyls .	
10 H 10 H	W.3	0CH <sub>3</sub>	н	×	н	OI3	סנוקטקטקיל \	CH <sub>2</sub>
-		0CH <sub>3</sub>	cu <sub>3</sub>	×	H		CH(CH <sup>2</sup> ) <sup>5</sup>	CH.
SO N	ᅄ	Ch.	- ,					

cont.
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	415	<b>a</b> 1	e <sup>2</sup>	۲,	R <sup>4</sup>	<b>8</b> 5	a <sup>6</sup> a <sup>7</sup>	R.O
_	<del>-</del>	<u> </u>	acı,			н	H -(CH <sub>2</sub> )4-	
	-		00h	H	N	×	# -{Di <sub>2</sub> )4-	
,	•		00H <sub>3</sub>	N	H	M	-(OI <sub>2</sub> )4*	
	-	- H	00kg	н	×	M	-(CI2)4-	×
•			004	*		H	H -0-{OI <sup>2</sup> }3-	
٥		N N	9CH <sub>2</sub>	. *	. #	M	H -0-(CI <sup>Z</sup> ) <sup>2</sup> -	
		×	oos,	H	H	н	-(CH <sup>2</sup> ) <sup>2</sup> -0-	
٥	11	M	0CH <sub>2</sub>	и		H	-{CH <sub>2</sub> } <sub>2</sub> -0-	
	_		00l <sub>3</sub>	*	×	×	H -OI-CH-CH-CH-	
0	-		0CH <sub>3</sub>	N			N -OH-CH-CH-CH-	
	-		OCH <sub>3</sub>	и	н	· •	-OI-OI-OI-	Ħ
			9CH <sub>3</sub>	×	W	×	-OI-OI-OI-OI-	Ħ
		M.	o( )	×	W	×	CH <sub>3</sub> 0CH <sub>3</sub>	оч <sub>3</sub>
ja	*	ĸ	٥٩	×			CH <sup>3</sup> 0CH <sup>3</sup>	cu3
			CH(OCH <sup>3</sup> ) <sup>5</sup>	n			CH3 OCH3	O13
•	-	-	3.5					COR E.

cent.

	16		R <sup>2</sup>	1,3	84	R <sup>S</sup>	26	g <sup>2</sup>	R <sup>B</sup>
_	R <sup>15</sup>	_R1						0CH <sub>3</sub>	ОŊ
Ø	×	18	CH(OCH <sup>3</sup> ) <sup>5</sup>	×		•		0CH <sub>3</sub>	Ol <sub>3</sub>
\$		×	040	н	н		-	0CH <sub>2</sub>	, O.
مو	H T	M	CH0	Ħ	H	×	Ol3	•	CH <sub>3</sub>
s	н	×	CH-CH-COOC <sub>2</sub> H <sub>5</sub>	М	H	М	ol3	0CH <sub>3</sub>	
50	×	H	OH-OH-COOC-HS	H	н	Ħ	CH3	OC)3	CN3
 s			۵۱ <sub>۲</sub> ۵۱۲ <sub>۲</sub> ۵۵۲ <sup>۲</sup> ۲۹	M	н	×	or <sup>3</sup>	OCH3	Dia
50	-		פולטולנוננילוול	×	н	×	· cH <sub>3</sub>	0C) <sup>3</sup>	Cr.)
-			Ol <sup>z</sup> Ol <sup>z</sup> Col(Col <sup>3</sup> ) <sup>5</sup>	н	×	*	c <sub>iO</sub>	OCH3	CI)
\$		-		н	н	×	CH3	0CH <sub>3</sub>	CH <sub>3</sub>
50	M	×	Or <sup>2</sup> Or <sup>2</sup> CON(Or <sup>2</sup> ) <sup>5</sup>	н	н	N	οij	0CH <sub>3</sub>	Oly
\$	×	ĸ	CH-CH-CH	и	н	×	CH <sub>3</sub>	OCH <sub>3</sub>	CH <sub>2</sub>
50	*	M	OI=OI-CII			H	οί,	0CH <sub>3</sub>	· OI <sub>2</sub>
5	×	H	DISCHIO	14			o+ <sub>3</sub>	0CH <sub>1</sub>	CH.
50		M	Ol <sup>2</sup> Ol <sup>2</sup> Ol	×	ж	ж		0CH <sub>3</sub>	01
\$	n	×	CH <sup>2</sup> CH <sup>2</sup> CH <sup>2</sup> CH	н	н	H	O+3	-	OI
so	16	×	Ol <sup>S</sup> Ol <sup>S</sup> Ol <sup>S</sup> ON	н	н	н	CH <sup>3</sup>	OCH <sub>3</sub>	
s		н	CHICHICOLOCOCHI	н	×	×	ᅄ	осн3	CH
50		м	CH <sup>2</sup> CH <sup>2</sup> CH <sup>2</sup> OCOCH <sup>3</sup>	н	Ħ	H	CH <sup>3</sup>	OCH3	CH
``			DIZCHZCHZII(CH3)2	н	н	×	ᅄ	осн	O
-				×	н	H	CI <sub>3</sub>	OCH3	O
50	M	н	01201201211(013)2				•		

	a <sup>15</sup>	<b>B</b> 1	<b>n²</b> ,	ره	84	45	16	1,	4
			OLOLOLUSCOC, N		4	н ,	OI,	och <sub>3</sub>	u,
,			OLOGOLIBEOC <sub>Z</sub> IL	×			cu	oci3	CH <sub>3</sub>
,	-		01-01-0001				c <sub>1</sub>	осн3	OI)
,	_		0-0000	×	H	*	013	OCH <sub>3</sub>	ᅄ
	-		01,01,0001	N			CH3	901 <sub>3</sub>	CH3
)	=	-	Or'Or'000r'				OI,	0CH <sub>3</sub>	CH <sub>3</sub>
		-	o⊷oı-(C)	•	H		DI <sub>3</sub>	ocu <sub>3</sub>	ᅄ
	_	-	a-a-(i)		M	*	Dij	оси <sub>3</sub>	. 043
		-	محمدی			H	CH <sub>3</sub>	0CH <sub>3</sub>	CH3
		#	مئيئن				043	0013	Or)
			رن کیمٹی	OL,		×	CH3	001201-012	CH <sub>3</sub>
		Oly To	-	ᅄ			O13	001201-012	013
ı	K	CH <sub>3</sub>	~ -(5)			H	CH <sub>3</sub>	OCH <sub>3</sub>	O <sub>3</sub>
	**		a,-⊙ ~ -⊙	-		H	013	9013	043
		<b>.</b>	oy(⊙)	-		ж .	04	8CH <sub>3</sub>	D13
	٠.	, <b>a</b>	<b>.</b> ***	-		n	Oly	OCH <sub>1</sub>	cu <sub>3</sub>
ı	M	*	0-(O) acu,ou, (O)	-		и	oi,	0CH <sub>3</sub>	013
	N		🔀				Oly	0CH <sub>2</sub>	01,
•			ecron (Q)	-			. •		

				-	<u>`</u> _		R6	g <sup>7</sup>	7.0
£1:	5	R <sup>1</sup>	g2	R3	14	R <sup>5</sup>	R*		
			001 <sub>2</sub> 01	×	×	×	o₁3	OCH <sup>3</sup>	CH)
		-	001 <sub>2</sub> CH	H	н.	H	, oi <sup>3</sup>	OCH3	C <sub>IC</sub> O
		_	סטילנטטב <sup>ל</sup> ווי	H	×	×	013	OCH3	C <sub>1</sub> 2
			001,000,14	H	ĸ		CH <sub>3</sub>	0CH <sub>3</sub>	C <sup>1</sup> 3
		-	• • •	×	×	H	c <sup>k</sup> 3	OCH <sub>3</sub>	ᅄ
×		*	ociforial *	. #	и.	×	OI <sub>3</sub>	0CH <sub>3</sub>	CH3
			BOITOITON	-	×	H	043	0CH)	OI;
H		И	001/01/0001-(0)	*	*	H	01,	OCH <sub>3</sub>	043
*		#	cortariocopy -(C)			и	OH.	0CH <sub>3</sub>	OI:
Ħ		×	OCIZOIZIBIZ			н	OH <sub>3</sub>	OCH <sub>2</sub>	OI.
		×	סטויטויאון איי	-	*		Ol <sub>3</sub>	OCH <sub>2</sub>	CH <sub>2</sub>
*		×	001 <sup>2</sup> 01 <sup>2</sup> INC001(01 <sup>2</sup> ) <sup>5</sup>		. ж	ж	o,	0CH <sub>3</sub>	CH
) ×		M	OOPDINGO(OI2)\$				οι,	· 0CH <sub>2</sub>	CH.
		M	oorteo -©	, <b>#</b>		*	CH <sub>3</sub>	OCH <sub>1</sub>	CH
ه ه		Ħ	∞ <sup>2</sup> co –⊘	11	-		CH2	осн <sub>3</sub>	O
, <b>x</b>	•	H	` <b>a</b> -∕ <b>©</b>	×	И		OI3	OCH <sub>3</sub>	OI
<b>o</b> 11	ł	H	∞-⊘> <u> </u>	н	N	# 	CH <sup>3</sup>	0CH <sub>2</sub>	CH
, н		H	co(cy <sup>2</sup> )30	ĸ	×	H**	•	-	0
ء م		H	co(Cor <sup>2</sup> ) <sup>2</sup> 0(C)	•	H	M	O13	OCH 3	
			$\leftarrow$	<b>a</b> .	14	N	OI)	OCH3	. 0

ALC: NAME: 1

<u>.</u>

	R15		2	<sub>2</sub> )	-,4	R5	26	R <sup>7</sup>	- 4
			<u>.</u>			*	DI,	001 <sub>3</sub>	CH3
50			~~	 (N <sub>3</sub>	H		OI,	OCH3	Ot3
5	Ħ		C000120120C13	on,	н	×	OI <sub>3</sub>	ocn <sub>3</sub>	CH3
50	H	-	cocos²(O) cocos²osòcos²	. Ol <sub>3</sub>		*	Olj	OCH <sub>3</sub>	013
•	H	•		. •			CH <sub>3</sub>	OCH <sub>3</sub>	CH <sub>3</sub>
0			യാം പ	о <sub>й</sub> ,		1	01	. оск <sub>3</sub>	. OI3
5	ĸ	×	Dr <sup>2</sup> OH	_		1	Oh	0CH <sub>3</sub>	ᅄ
<b>50</b>	Ħ	*	DYDN	ch <sub>3</sub> ~	11		013	001 <sub>3</sub>	οŋ
\$	m		or <sup>1</sup> 000{(O)	O <sup>1</sup> 3	- *		Ol <sub>3</sub>	0CH <sub>3</sub> •	ο'n
54	11	ĸ	01/00 -(O)	ο <sub>3</sub>	»	×	CH <sub>3</sub>	001201-012	CH3
\$	Ħ	ĸ	coco,	O <sup>4</sup> 3	 p	*	Oly	001 <sub>2</sub> 01-01 <sub>2</sub>	O' <sub>3</sub>
50	*	1	COOCH	ы Сы <sub>р</sub>			013	ecs <sub>3</sub>	Ol <sub>3</sub>
\$	•	H	DI <sub>Z</sub> CI <sub>Z</sub> OCII <sub>3</sub>			×	Ol <sub>3</sub>	0CH <sub>3</sub>	CH <sub>3</sub>
50	Ħ	*	סילטולסטו	*	-		oi,	001501-015	CN <sub>2</sub>
\$	*	*	01(013)2	**	-	-	Ol <sub>3</sub> .	•	ОЧ,
50	Ħ	H	GI(GI <sup>3</sup> ) <sup>5</sup>		-	- H	0/3	001,01-01,	OI,
\$	H	*	c(cx3)3	*	-		οı	001201-012	CH.
\$0	Ħ		c(cx <sup>3</sup> ) <sup>3</sup>				CK <sub>3</sub>	6013	OI,
5	M	OI,	OCH <sub>3</sub>	CH3	• .		,	•	

cont.

	R <sup>15</sup>	<u></u>	2	R <sup>3</sup>	R <sup>4</sup>	R <sup>S</sup>	26	ري	
_			001 <sub>3</sub>	CH3	м	×	CH3	0013	, Or <sup>3</sup>
50		O'3	осн <sub>а</sub>	о <sub>3</sub>	ĸ	И	CH3	CH <sub>3</sub>	H
\$	M	ο' <sub>3</sub>	•	OI <sub>3</sub>	×	×	CH3	OI <sub>3</sub>	H
50	H	o's ~	001 <sub>3</sub>	CH <sub>3</sub>	н	н	OI <sub>3</sub>	9CH <sub>3</sub>	CH3
<b>S</b>	<b>H</b>	о <sub>3</sub> ~	סטו <sup>ל</sup> טולסטו <sup>3</sup>	CH <sub>3</sub>	H	н	CH <sub>3</sub>	OCH <sub>3</sub>	ᅄ
50	*	ο <sub>3</sub>	מטו <sup>ב</sup> טר <sup>ב</sup> טנא <sup>3</sup>	οι <sub>3</sub>	н	ĸ	H	CK3	OI3
\$	и	O43	מסולטולטטול	οι <sub>1</sub>	и	н	×	CH <sub>3</sub>	CH3
50	M	O'S	001 <sup>2</sup> 01 <sup>2</sup> 001 <sup>3</sup>	си <sub>3</sub>	и	H	CH <sub>3</sub>	OCH3	CH <sup>3</sup>
5	<b>N</b>	οl <sub>3</sub>	COCK3	οι3 .	×	ж .	CH <sub>3</sub>	OCK <sub>3</sub>	c <sub>i</sub> 3
50		Ol <sub>3</sub>	coci <sup>2</sup>	CH <sub>3</sub>	н	н	CN <sub>3</sub>	H	CH3
\$		O13	COCH <sup>3</sup>	ск,	н	н	CH <sub>3</sub>	ж	CHO
50	¥	CH <sub>3</sub>	COCH <sup>3</sup>	CH3	×	н	CH3	осн <sub>3</sub>	ĊH <sub>3</sub>
\$	<b>N</b>	CN <sub>3</sub>	coc <sub>z</sub> u <sub>s</sub>	CH <sub>3</sub>	ж	н.	CH <sub>3</sub>	осн <sub>3</sub>	OI3
50	H	CH3	COC2H2	CH3	и	н	CH3	осн3	CH3
\$	CH.3	CH <sup>3</sup>	си3	CH3	H	н	CH <sub>3</sub>	OCH <sub>3</sub>	CH <sub>3</sub>
20	CH <sub>3</sub>	CH <sub>3</sub>	CH <sup>3</sup>	си <sub>з</sub>	н	ж	OH <sub>3</sub>	CH3	CH3
\$	н	CH <sub>3</sub>	CH <sub>3</sub>	•		н	OI <sub>3</sub>	CH <sub>3</sub>	CH <sub>2</sub>
50	N	CH <sub>3</sub>	сн <sub>3</sub>	CH3			CH3	OCH <sub>2</sub>	CH <sub>3</sub>
\$	×	CH3	C2H5	CH3	••	.,	•		cont.

cont.

cent.

	£15	<u>,1</u>	g <sup>2</sup>	. 83	14	25	16	A <sup>7</sup>	R <sup>8</sup>
<u>.</u>				O13		и	OI)	0CH <sub>3</sub>	CH3
		o₁ ~	ር Ju	οί,	×	×	CH3	оси <sub>3</sub>	М
<b>.</b>		α <sub>3</sub> ~	c <sub>Z</sub> u <sub>s</sub>	CI <sub>3</sub>	*	*	OI3	<sup>001</sup> 3	Ħ
		о <sub>3</sub> ~	Chr.	013	н	M	OI3	осн <sub>3</sub>	Ch2
<b>.</b>		O <sub>2</sub>	DI(DI <sup>3</sup> ) <sup>5</sup> DI(DI <sup>3</sup> ) <sup>5</sup>	Ol <sub>3</sub>	×	, <b>N</b>	CH <sub>3</sub>	осн <sub>3</sub>	CH <sub>3</sub>
<b>.</b>		O'3		ο. <sub>1</sub>		и	CH <sub>3</sub>	CH <sub>3</sub>	CH3
		ol <sub>3</sub>	Or(Or <sup>2</sup> ) <sup>5</sup>	CN <sub>3</sub>	н	×	01,	CH <sub>3</sub>	OI,
0	H	ο <sub>3</sub>	01(01 <sub>3</sub> ) <sub>2</sub>	; Ol <sub>3</sub>		×	CH <sub>3</sub>	OCH <sub>3</sub>	CH)
		Ol3	αοι <sup>γ</sup> -(∴)	013		H	CH <sub>3</sub>	оси <sub>з</sub> .	cu <sup>3</sup>
0	•	CH <sub>3</sub>	- <b>-</b> .	0CH <sub>2</sub>	H	и .	OI3	0CH <sub>3</sub>	CH <sub>3</sub>
•	M	0CH <sub>3</sub>	Br Di	0CH <sub>3</sub>	ж	×	OK <sub>3</sub>	0013	CH3
540	*	905	Br .	OCH <sub>3</sub>	. н	N	Ol <sub>3</sub>	CH <sub>3</sub>	×
	×	och,	Br .	0CH <sub>3</sub>	н'	н	013	OI <sub>3</sub>	, 1
50	×	001 <sub>3</sub>	br ~	•			οij	OCH <sub>3</sub>	CH3
•	×	chil	CII	C <sub>Z</sub> N <sub>S</sub>		н	01,	OCH <sub>3</sub>	ᅄ
so	N	وكهلا	CR	C <sub>2</sub> H <sub>5</sub>	и .	×	CH <sub>3</sub>	0C <sub>2</sub> N <sub>5</sub>	Oi <sub>3</sub>
S		CZNS	CI.	CzNs	 H	н	οι,	OC <sub>2</sub> N <sub>2</sub>	CI <sub>3</sub>
50	H	وكالح	OI .	CZHS		н	ο,	осн <sub>3</sub>	CH3
\$	M	O13	осн <sub>3</sub>	CH <sup>3</sup>	C <sup>1</sup> 3	-	- 1	•	•

cont.

				R <sup>3</sup>	14	R5	A <sup>6</sup>	R <sup>7</sup>	gå.
	R <sup>15</sup>	8 <sup>1</sup>	<u>r,</u>				O13	0CH <sub>1</sub>	043
0	H	OL3	0CH <sub>3</sub>	CH3	<sup>O1</sup> 3		-	00H <sub>1</sub>	oi,
	×	013	OCH <sub>3</sub>	×	OI3	H	CH <sup>3</sup>	-	CH <sub>3</sub>
٥	×	043	001 <sub>3</sub>	ĸ	CI(3	K	O13	0CH <sub>3</sub>	OI <sub>3</sub>
,	*	ci .	001 <sub>3</sub>	Ħ	OCH <sub>3</sub>	Ħ	CH3	OCH3	_
٥		¢1	001 <sub>3</sub>	H	OCH <sub>3</sub>	H	CH3	OCH 3	CH <sub>3</sub>
		cı	c1	C3	Ħ	M	CHO	0CH <sub>3</sub>	Ol3
•	H		c)	c1	м	И	CH <sub>3</sub>	OCH <sub>3</sub>	OI3
0	×	C)		c1		H	CH3	OCH-CH-CH2	OI3
	Ħ	C)	C1	c)	н	ĸ	он,	OCH <sup>2</sup> CH-CH <sup>2</sup>	OI3
٥	×	C1	C1		cı	н	OH <sub>3</sub>	OCH <sub>2</sub>	CK3
	Ħ	C1	C1	<b>C1</b>	61	 H	D13	0CH <sub>3</sub>	CN <sub>3</sub>
50	Ħ	cı	C1	C1		и	013	0CH2CH-CH2	013
s	H	E1	C3	¢1	c1		CH <sub>3</sub>	OCH <sub>2</sub> CH-CH <sub>2</sub>	n,
50	M	c1	c1	C)	C1	H	-	DOH <sub>3</sub>	о,
s		OCH3	Br	н	OCH <sup>3</sup>	M	Ol <sub>3</sub>	-	CH <sub>3</sub>
- 50	и	ocu <sub>3</sub>	lr .	н	OCH 3	H	CH3	OCH 3	-
~ s		OCH <sub>2</sub>	C1	¢1	OC <sub>2</sub> H <sub>5</sub>	H	CH3	OCH.)	CH <sub>2</sub>
		OCH <sub>3</sub>	c1	C1	oc <sub>z</sub> n <sub>s</sub>	H	CH3	осн <sub>3</sub>	CH
50	H	•	c)	cı	OC <sub>Z</sub> H <sub>S</sub>	н	CH3	CH3	H
\$	×	9013	.,	•				_	ent.

cont.

	a <sup>15</sup>	21 -	2	٠, ۲	A4	25	8.6	R <sup>7</sup>	م.
	8			<u> </u>	oc <sub>z</sub> v <sub>s</sub>		٥١,	CN <sub>3</sub>	M
0		oor <sup>2</sup>	ต		CH <sub>3</sub>		OI,	OCH <sub>3</sub>	OI3
•	Ħ	യവു	Oly	O13	Oly	*	CH,	0013	CN <sub>3</sub>
	×	cock3 .	ο <sub>3</sub>	O13	ei		OI <sub>2</sub>	001,	OI3
		F 75.5	a	, и	=	-	Ol <sub>3</sub>	OCH <sub>3</sub>	· cu <sub>3</sub>
0		F	CI	, N	C1	•	OI <sub>3</sub>	0CH <sub>2</sub>	CH <sub>2</sub>
	M	C1	Ormon <sup>2</sup>	c1	×	H	. •	00l <sub>3</sub>	OI,
۰	ĸ	ព	OLCOOO13	C1	. #	Ħ	Oly C	•	οί
	M	ព	ou <sub>z</sub> os	C1	M	K	, oi <sub>3</sub>	0CH <sub>3</sub>	ος
•		C1	ou, ca	cı	. #	я .	CN3	001 <sub>3</sub>	•
•	в.	-01-01	a-a-	oi-a	1-01-01-	*	Ol3	0CH <sub>3</sub>	Ol3
i		#	m )		i	H	cH3	00H <sub>3</sub>	cx3
			α <sub>3</sub> > ¬				· 01 <sub>1</sub>	0CH <sub>3</sub>	о'n
0	Ħ	# 1813. 1				H	OK <sub>3</sub>	0CH <sub>3</sub>	. Olg
0	×	и .			-	<u></u>	013	001,	o۱
	*	*	-∞	o-	H.		o <sub>3</sub>	оси <sub>з</sub>	α <sub>3</sub>
•	-	-		4					cont

cont.

LUII C.		•						وم
	R <sup>15</sup>	g <sup>1</sup> g <sup>2</sup>	R3	R <sup>4</sup>	R <sup>5</sup>	g <sup>6</sup>	R <sup>7</sup>	
_			-00120-	N	H	013	0CH <sub>3</sub>	CH3
0	×		-00/20-	H	N	013	c <sub>IO</sub> 3	CH3
; ;0	H		-00/o-	H	ĸ	cH3	cH <sup>3</sup>	CH3
5	18	H	Q.	×	H	O13	0CH <sub>3</sub>	CH <sup>3</sup>
50		a a	<b>`</b> \$	, **	н	OI3	001 <sub>3</sub>	DH.
	-	-01-01-01-11-	н	м	M	CH <sub>3</sub>	OCH <sup>3</sup>	CH3
5	<b>X</b>	-OI-OI-OI-II-	н	. н	н	, CH <sub>3</sub>	OCH <sub>3</sub>	CH <sup>3</sup>
50		•		н	H	OI3	0CH <sub>3</sub>	CH <sub>3</sub>
5	н	-OI-OI-OI-OI-		H	H	CH <sub>3</sub>	0CH <sub>3</sub>	CH3
50	*	-01-01-01-01-	O-OI-OI-OI-	<b>H</b>	н	CH <sub>3</sub>	OCH <sub>3</sub>	CH3
S	H	. #	-O+O+-O!-	н	×	DI <sub>3</sub>	OCH <sub>3</sub>	CH3
50	×	И	-O-O-O-O-	н	×	OI,	осн <sub>3</sub>	CH <sub>3</sub>
S	H	-01/01/01/01	3	• и	*	CH <sub>3</sub>	осн	cu <sub>3</sub>
50	н	-01201201201	2° H			_		CH3
5	H	DO13	012D12D12	C1	н	CH3	0CH <sub>3</sub>	-
so		0CH <sub>3</sub>	-01,01,01,-	C1	H	CH <sub>3</sub>	0CH <sub>3</sub>	
5	н	οοι <sub>3</sub>	- میرمارمار-	C1	и	Ol3	oc <sub>z</sub> n <sub>s</sub>	CH <sup>3</sup>
•	••	3	. • •					con

l.	R <sup>25</sup>	81	R <sup>2</sup>	g <sup>3</sup>	14	g\$	10	R <sup>J</sup>	/ g8,
8		<b>60</b> 5		-01,01,01,-	<b>C1</b>	И	013	oc <sub>z</sub> n <sub>s</sub>	. OI <sup>3</sup>
		- 0	1 - 01 - 0	1 • C • DYDIZ •			CH <sub>3</sub>	0CH <sub>3</sub>	Or <sup>2</sup>
٥	*	- 0	ı - C1 - C	x = c - OL <sub>2</sub> OL <sub>2</sub> -			013	0013	c <sup>ب</sup>
•	., .			© <u>,</u> ·			OI3	оск <sub>а</sub> .	α,
•	w	ж	••	<b></b> _	×	×	013	`0CH <sub>3</sub>	α <sub>3</sub>
	×			-001-0-	×	C0,CH <sub>3</sub>	013	001 <sub>3</sub> ;	CP3
•	16	H		-001,0-	*	00,013	013	0CH3	OL <sub>3</sub>
				-001,0-	H	CO2C2HS	OI <sub>3</sub>	9CH <sub>3</sub>	OI3
•				-001,0-	×	CO <sub>2</sub> C <sub>2</sub> H <sub>2</sub>	CH3	0CN3	cx <sup>3</sup>
,		H	•	-00150-	*	co <sub>2</sub> c(O( <sub>3</sub> ) <sub>3</sub>	οij	0013	CH <sub>3</sub>
•		N		-0040-	H	ω <sup>2</sup> ε(cn <sup>3</sup> ) <sup>3</sup> (	CI3	OCH3	ot <sub>2</sub>
				-00120-	. H	00,01,-(C)	" CN <sub>3</sub>	0CH <sub>3</sub>	OI3
,	H	N		-001 <sub>2</sub> 0-	H	صمت ⊘	013	0CH <sub>3</sub>	013
	#	×		-0CH <sub>2</sub> 0-	H	<b>∞</b> -⊚_	CH <sub>3</sub>	0CH <sub>3</sub>	CH3
•	н	Ħ		-0CH <sub>Z</sub> 0-		<b>∞ ⊘</b>	C×3	OCH <sup>3</sup>	CV <sub>3</sub>
	×	И		-00120-	н	comi	OI <sub>3</sub>	0013	CN <sub>3</sub>

cent.

	R <sup>15</sup>	<b>B</b> <sup>3</sup>	R <sup>2</sup>		R3	R <sup>4</sup>	R5 -	R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>
				-001,0-		H	COMIZ	OI3	0CH <sub>3</sub>	CK3
	_	- 1		-00120-		H	COUNTY	CH3	0CH <sub>3</sub>	CH3
	-			-00/20-			CONICZIIS	ыş	0CH <sup>3</sup>	ᅄ
				-00/0-		н '	COMOLY-(C	) o <sub>3</sub>	OCH 3	CH3
				-001,0-		×	сонюц{(	∑) o₁,	9CH <sub>3</sub>	cx³ .
-		»		-0050-		н	COMM(CO)	O13	оси <sub>3</sub>	CH3
٥	 H	 M		-001 <sub>2</sub> 0-		×	· coset —((())	CN3	0CH <sub>3</sub>	CH3
•	 M	 H		-001-0-		н	con(CH <sub>3</sub> )2	013	осн <sub>э</sub> .	CII3
.o		 M		-0040-		н	CON(CH <sub>3</sub> ) <sub>2</sub>	ᅄ	0CH <sub>3</sub>	O13
~ i		οι,	OI <sub>3</sub>	•	CH <sub>3</sub>		н	_ CH <sub>3</sub>	001,01,001	O13
50	ж	οι <sub>3</sub>	ο,		OI,	n	<b>, w</b>	OI,	00H2CH200H3	cı,
•	н	H .	OCH.		» T	н	Ħ	-0	H+CH-0-	Ħ
	 M	 H	001		н	н	M	-0	H-CH-0-	H
5		H	0013		H	. *	н	н	-0-OI-	CH-
50	и.		OCH3	•	н	н	H	И	-ò-OI-	CH-
 د	" *	H	0013		н	H	H	-0	H-CH-HH-	H
50		ĸ	OCH <sub>3</sub>		N	н	н	-0	N=CH- <del>INI</del> -	H
30 S			OCH <sub>3</sub>		×	н		н	-W-O	<b>-O-</b> .

	a15	۲)	e <sup>2</sup>	R <sup>3</sup>	**	<b>2</b> 5	R <sup>&amp;</sup>	R <sup>7</sup>	• •
,			9CH <sub>3</sub>	×	и	×	×	•	H-CH+CH+
			0CH <sub>2</sub>	H	N		-01-	OI-8(OI <sub>3</sub> )-	×
,	_		ock <sub>3</sub>		×	*	-01-	OI-11(DI <sub>3</sub> )-	H
-	-		00L	N		и	н	-A(C	*3)-CH+CH-
	-	 B	, 001 <sup>3</sup>	×	*	N	н	- un , -11(	012)-01-01-
•	-	 O4	Orcioi	CH <sub>3</sub>	M		o۱	0013	CH <sub>3</sub>
3		. Dr.	Or'C+Ot	01,	Ħ	H	Oi,	0CH <sub>3</sub>	cu <sub>3</sub>
•		-73	محمدها الم	H ,		×	01,	0CH <sub>3</sub>	cr <sup>3</sup>
	-		المراحية	×		H	_ CH <sub>3</sub>	0CH <sub>3</sub>	CH <sub>3</sub>
•	-	-	corororo	×	×		CH <sub>3</sub>	0CH <sub>3</sub>	C <sub>IC</sub>
•	-	-	معاماتهای			N	CH <sub>3</sub>	9CH <sub>3</sub>	OI3
•		Ol <sub>2</sub>	6(01 <sup>2</sup> ) <sup>2</sup> 01 <sup>3</sup>	OI <sub>3</sub>		×	œ	oci <sub>3</sub>	cu <sup>3</sup>
•		01 <sub>2</sub>	6(ci <sup>2</sup> ) <sup>2</sup> ci <sup>3</sup>	οι,	H	*	013	OCH <sub>3</sub>	013
	-	<b>3</b>	cyts	×	*	×	CH <sub>3</sub>	001201-012	OI,
•	-	*	GR.	H	Ħ	<b>*</b> .	v 013	001201-012	OI <sub>3</sub>
•	-	_	001) 23	*	*	<b>∞-</b> ⊘	DI <sub>3</sub>	0CH <sub>3</sub>	01)
,	-	-	<u>-</u>	001 <sub>3</sub>		∞-⟨Ō⟩	OI <sub>3</sub>	0CH <sub>3</sub>	CH <sub>3</sub>
, 10	-	-	0CH <sub>3</sub>	н		ω-(Ō̄)	043	0013	CH <sub>2</sub>
_	-	-							COR

	R <sup>15</sup>	11	r <sup>2</sup>	<b>8</b> 3	1,4	g.S	R <sup>6</sup>	ę?	gå
_	<del>-</del> -	<u> </u>		0CH <sub>2</sub>	* H	u-(⊙) ÷	۵,	OCH3	CH3
50 S		-	 Ск <sub>а</sub>	OIZOCO (C	S) #		CH3	ocs <sup>3</sup>	OI3
			or <sup>2</sup> 000 -{○}	· 01,		<b>∞</b> · <b>⊘</b>	OI3	оси <sub>3</sub> ·	о,
		•	-001 <sub>2</sub> 0-		н	coczns	013	0CH <sub>3</sub>	cu <sub>3</sub>
i io		-	-001-0-		*	COCTIL	СIO	ocn <sup>3</sup>	ο <sub>b</sub> , ·
~ 10	<b>N</b>	*	οι <sub>3</sub>	clo	*	COOCH <sup>3</sup>	91,	0CH <sub>3</sub>	Crl3
s		-oc*	<u> </u>	*	H	N ·	œ3	оси <sub>3</sub>	013
SO.		-00	× <u>,</u>	N	• и	н	CH3	0013	cu3
- s			SOI3	н	н	H	CH3	ooi <sub>3</sub>	o,
s			CH(CH <sub>3</sub> ) <sub>2</sub>	Ħ	H	н ,	CH3	осн <sub>2</sub> -<	сı,3
50		ĸ	СH(СН <sup>3</sup> ) <sup>5</sup>	H	• н	н	CH3	0CH <sup>2</sup> -{0}	cu3
s			CH <sup>2</sup> CH <sup>2</sup> COCH <sup>3</sup>	н	M	×	C×3	OCH <sup>2</sup> CH-CH <sup>2</sup>	CH3
90			снуснусосну	н	×	H	CH3	OCH <sup>2</sup> CH-CH <sup>2</sup>	CH3
50	,. M		CH <sub>3</sub>	сиз	H	COOC(CH <sup>3</sup> ).	CH3	OCH <sub>3</sub>	CH3

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Table 1 cont.

	•	15	11		g2		R <sup>3</sup>	R <sup>4</sup>	A.5	R <sup>6</sup>	a,7	- R	
-					City		CH <sub>2</sub>	*	con(CH <sub>2</sub> ) <sub>2</sub>	CH)	OCH <sub>3</sub>	a	×3
•	•		_		01,		OI,	#	con(CH <sub>3</sub> ) <sub>2</sub>	CH3	OCH <sub>3</sub>	C	×3
	_			4,	, 1		n	×	H	CH <sub>3</sub>	*DOI*CII-CII*	. C	×
			-				H	H		CH <sub>3</sub>	001201-012	C	<b>~</b> >
	د .	- 1-	OL.	<u>-</u>	CIL.	::	OL	H	H	O13	OI3	×	ţ
•			С		OL.		οų	Ħ		CH <sub>3</sub>	CH <sub>3</sub>	. 1	i .
		•			CI.	j.	OL	Ħ		M	CH <sub>3</sub>	Ĉ	×3
				: <u>;</u>	OL.	**	οų				CH <sub>3</sub>	C	×3
~	′ ]	77.7. 1 = 3	ΩL.		CU.	·-,	CUL	M		CH3		. , 0	<sup>24</sup> 3
-					OL.	Ĭ.	OL,	. #	N	CH3	ĸ		<sup>(2)</sup> 3
	_		. a.		OL.		×	. 013		043	CN3	1	•
		,,	. TJ	,	CIL.		M	CH <sub>3</sub>	H	CL)	CH <sub>3</sub>	. 1	4
7			െ		<b>a</b>	.:	CH <sub>2</sub>	×		CH <sub>3</sub>	oc <sub>2</sub> n <sub>5</sub>		<sub>01</sub> 3
	ं		OL.		CIA		CII <sub>1</sub>		. ж	cu <sub>3</sub>	oc <sub>z</sub> n <sub>s</sub>	(	Cr.
9		,			COOCH	• • • • • • • • • • • • • • • • • • • •	CH <sub>3</sub>	H			OCH <sub>3</sub>	(	ديالح
•						-cizoizoiz	• .	H	H	ᅄ	OCH <sub>3</sub>	•	CI(3

Table 1 cont.

	R <sup>15</sup>	R1	2	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>
<u>.</u> so	<u>.</u>		-CH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub>	•	н	н	CH3	осн3	CH3
50 50		 M	OCH <sub>3</sub>	н	н	Ħ	-CH <sub>2</sub>	сн <sub>2</sub> сн <sub>2</sub> 0-	. N
بد دو	H	 H	OCH <sub>3</sub>	H	×	M	M	-001201	•
بح د	N	W.	SOCH	H	11	ж <sub>.</sub>	CH <sub>3</sub>	OCH <sup>3</sup>	CH <sub>3</sub>
50	 M	16	· · · · SOCH <sub>2</sub> · · · · ·	н	н	M	CH3	OCH <sub>3</sub>	CH <sub>3</sub>
30 S	H	<b>x</b>	CH <sub>3</sub>	CH <sub>3</sub>	М	H	CH3	-0CH2-	CH <sup>3</sup>
50	н	×	СН <sub>3</sub> .	CH <sub>3</sub>	н	н	CH3	-0CH <sub>2</sub> -	СНЗ
ىد د			H-CH-CH-CH-	-	-сн-сн-	H	CH <sub>3</sub>	0CH <sup>2</sup>	CH3
20 20	×	N.	NO <sub>2</sub>	ж	н	н	CH3	осн <sub>3</sub>	CH3
\$	н	H	σ,	н	н	H	CH3	∞H <sup>2</sup> ✓	CH3
50	н	, н	σ,	H	н	Ж	CH <sub>3</sub>	осн <sub>2</sub>	CH3
5	 H	×	CH <sub>2</sub> CH <sub>2</sub> COOC <sub>2</sub> H <sub>5</sub>	н	н	н	CH3	осн <sub>3</sub>	CH3
sa sa	×	ĸ	осн3	н	н	<sub>0</sub> /C-0C(CH <sup>3</sup> ) <sup>3</sup>	CH <sub>3</sub>	осн3	CH3
50	×	n	CH3	CH <sub>3</sub>	н	H	H	осн3	с <sub>2</sub> н

The invention takes into consideration that compounds that structurally deviate from the formula I, after administration to a living organism may be transformed to a compound of formula I and in this structural form exert their effect. Such compounds structurally deviating from compounds of the formula I, are included in the scope of the invention.

Likewise, certain compounds of formula I may be metabolized into other compounds of formula I to before exerting their effect. Compounds of the invention wherein X is S are thus believed to exert their antisecretory and cytoprotective activities after metabolism to compounds wherein X is SO and compounds of the invention wherein R<sup>5</sup> is R<sup>14</sup>CO are believed to exert antisecretory and cytoprotective activity after metabolism to compounds wherein R<sup>5</sup> is

H. These considerations are also a further aspect of the invention.

Further, it is believed that all compounds of
20 formula I wherein X is SO after administration to a
living organism, exert their antisecretory and cytoprotective effects after metabolic or pure chemical
transformation to another, reactive species. Accordingly, the same is true also for the compounds of

25 formula I wherein X is S, but via initial transformation to the corresponding compounds of formula I wherein X is SO. These considerations as well as such reactive species per se are included within the scope of the present invention.

30 Preparation

Compounds of formula I above may be prepared according to the following methods:

a) Oxidizing a compound of the formula l,

wherein X is S and R<sup>15</sup>, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> have the meanings given, to give a compound of the same formula I wherein X is SO. This oxidation may 5 be carried out by using an oxidizing agent selected from the group consisting of nitric acid, hydrogen peroxide, peracids, peresters, ozone, dinitrogentetraoxide, iodosobenzene, N-halosuccinimide, I-chlorobenzotriazole, t-butylhypochlorite, diazabicyclo-

10 [2,2,2] - octane bromine complex, sodium metaperiodate, selenium dioxide, manganese dioxide, chromic acid, cericammonium nitrate, bromine, chlorine, and sulfuryl chloride. The oxidation usually takes place in a solvent wherein the oxidizing agent is present in some excess in relation to the product to be oxidized.

The oxidation may also be carried out enzymatically by using an oxidating anzyme or microbiotically by using a suitable microorganism.

20 b) Reacting a compound of the formula

with a compound of the formula

in which formulas R<sup>15</sup>, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> are as defined previously and wherein one of Z<sup>1</sup> and Z<sup>2</sup> is SH and the other is a leaving group, gives a compound of the formula I wherein X is S.

Examples of leaving groups Z<sup>1</sup> and Z<sup>2</sup> in the compounds II and III are halogens, preferably chlorine, bromine or iodinem acyloxy radicals, for example residues of strong organic sulfonic acids, for instance of an arylsufonic acid, for example tosyloxy or an alkylsulfonic acid, for example mesyloxy, alkylmercapto groups, for example methylmercapto, alkylsulfinyl groups, for example methylsulfinyl and the like.

Thus, Z<sup>1</sup> or Z<sup>2</sup> when designating leaving groups may be a reactive esterified hydroxy group. The esterification may be carried out with an organic acid or with an inorganic a .id such as HCl, HBr or H<sub>2</sub>SO<sub>4</sub>.

The reaction of a compound of formula II above
40 with a compound of formula III is conveniently carried
out in the presence of a suitable solvent that is inert
under the reaction conditions utilized as described
hereinafter. The reaction may further be carried out in
the presence of a suitable base. Suitable, bases
45 include, for example, inorganic bases such as sodium

or potassium hydroxide, sodium or potassium alkoxide, sodium or potassium hydride and the like, organic bases such as tertiary amines, for example triethylamine and the like.

Suitable solvents for the above described reaction include, for example, alcohols, preferably lower alkanols such as methanol and ethanol mixtures of such alcohols with water, ethers, such as tetrahydrofuran, halogenated hydrocarbons, such as methylene chloride. Aprotic solvents such as ethers and halogenated carbons are necessary in the case of sodium and potassium hydride.

The reaction of the compounds of formulas II and III may be carried out at a temperature between the ambient temperature and the boiling temperature of the reaction mixture. It is preferred to carry out the reaction, however, at a temperature at or close to the boiling point of the reaction mixture for the preparation of a compound of the formula I wherein R<sup>5</sup> is H.

e) Esterification of a compound of the formula

 $R^{\frac{1}{2}} \xrightarrow{R^{\frac{1}{2}}} R^{\frac{1}{2}} \xrightarrow{R^{\frac{1}{2}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}}} \sqrt{\frac{1}{R^{\frac{1}{2}}}}} \sqrt{\frac{1}{R^{\frac{1$ 

wherein R<sup>15</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> are as defined above and Y<sup>1</sup>, Y<sup>2</sup>, Y<sup>3</sup> and Y<sup>4</sup> represent either R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> according to the above definition, respectively, or the groups (Z) <sub>n</sub>-A-COOH, COOH and (Z)<sub>n</sub>-A-OH, whereby Z, n and A are as defined above, by reaction with the appropriate alcohol R<sup>9</sup>OH, R<sup>10</sup>OH or carboxylic acid R<sup>10</sup>COOH, respectively, to the formation of a compound of formula I containing a radical R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and/or R<sup>4</sup> which is either of the ester groups (Z)<sub>n</sub>-A-COOR<sup>9</sup>, COOR<sup>10</sup> or (Z)<sub>n</sub>-A-OCOR<sup>10</sup>.

The esterfication is carried out as an ordinary esterfication, in the presence of an acid catalyst such as sulfuric acid, hydrochloric acid and p-toluenesulphonic acid and, if necessary, in the presence of an inert solvent such as toluene.

d) Acylation of a compound of the formula

wherein  $R^{15}$ , X,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ ,  $R^7$  and  $R^8$  are as defined above, by reaction with an appropriate acylating agent ( $R^{14}CO$ )<sub>2</sub>O,  $R^{14}COX^1$ , whereby  $X^1$  is a leaving group such as C1,  $N_3$  and p-nitrophenoxy,  $R^4NCO$ , whereby  $R^8$  is defined by the relation  $R^8NH$  equals  $R^{14}$ , provided that  $R^8$  is K when  $R^{14}$  is amino, to the formation of a compound of formula I wherein  $R^5$  is  $R^{14}CO$  as defined above.

The acylation is preferably carried out in the presence of a base such as triethylamine, K<sub>2</sub>CO<sub>3</sub> and NaOH and with a solvent such as tetrahydrofuran, acetonitrile and water. Normally, if the benzimidazole moiety is asymetrically substituted, both the N(1)-

and the N(3)-acyl derivatives are obtained, and therefore, if necessary, the two components have to be separated. This may be done by recrystallizations or by extractive or chromatographic techniques.

e) Hydrolyzing a compound of the formula

wherein X,  $R^{15}$ ,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^6$ ,  $R^7$  and  $R^8$  are as defined above and Z3 is a suitable N-protecting group such as alkanoyl, carboalkoxy and trimethylsilvl, to the formation of a compound of the formula I wherein 10 R<sup>5</sup> is H.

The alkanoyl group in Z<sup>3</sup> can have 1-6 carbon atoms and the carboalkoxy group 2-6 carbon atoms. The hydrolysis may be performed in alkaline solution or in . acidic solution, the latter mainly for compounds 15 wherein X is S:

whereafter the compound of the formula I obtained if desired, when X is -S-, is converted to a physiologically acceptable salt or oxidized to form a compound of the formula I wherein X is -SO-.

Depending on the process conditions and the starting materials, the end products of the formula l wherein X is S is obtained either as the free base or as a salt. The end products of the formula I wherein X is, -SO- are obtained as the free base. Both the free base

25 and the salts of these end products are included within the scope of the invention. Thus, basic, neutral or mixed salts may be obtained as well as hemi, mono, sesqui or polyhydrates. Acid addition salts of the new sulficides may in a manner known per se be

30 transformed into free base using basic agents such as alkali or by ion exchange. The free bases of the selfides obtained may also form salts with organic or inorganic acids. In the preparation of acid addition salts preferably such acids are used which form 35 suitable therapeutically acceptable salts.

Examples of such acids are hydrohalogen acids, sulfonic acid, phosphoric acid, nitric acid, and perchloric acid; aliphatic, alicyclic, aromatic or heterocyclic carboxyl or sulfonic acids, such as formic acid, 40 acetic acid, propionic acid, succinic acid, glycolic acid, lactic acid, malic acid, tartaric acid, citric acid, ascorbic acid, maleic acid, hydroxymaleic acid, pyruvic acid, phenylacetic acid, benzoic acid, p-aminobenzoic acid, p - hydroxybenzoic acid, salicyclic 45 acid or p-aminosalicylic acid, ambonic acid, methanesulfonic acid, ethanesulfonic acid, hydroxyethanesulfonic acid, ethylenesulfonic acid, halogenbenzenesulfonic acid, toluenesulfonic acid, naphtylsulfoni - acid or sulfanilic acids, methionine, 50 tryptophane, lysing or arginine.

These or other salts of the new sulfide compounds, as e.g. picrates, may serve as purifying agents of the free bases obtained. Salts of the bases may be formed, separated from solution, and then the free 55 base can be recovered in higher purity from a new

salt solution. Racemates obtained can be separated according to

known methods, e.g. recrystallization from an optically active solvent, use of microorganisms, reactions 60 with optically active acids forming diastereomeric salts which can be separated, (e.g. separation based on different solubilities of the diastereomers), acylation of the benzimidazole nitrogen ( $R^5 = H$ ) or another nitrogen or oxygen atom in a substituent by an 65 optically active activated carboxylic acid (e.g. acid chloride), followed by chromatographic separation and deacylation.

Suitable optically active acids for salt formation are the L- and D-forms of tartaric acid, di - o - tolyl - tartaric 70 acid, malic acid, mandelic acid, camphorsulfonic acid or quinic acid, and for acylation O - methylmandelic acid. Preferably the more active part of the two antipodes is isolated.

In the case of diastereomeric mixtures (racemate 75 mixtures) these may be separated into stereoisomeric (diastereomeric) pure racemates by means of chromatography or fractional crystalliza-

The starting materials utilized in the processes a and c-e are obtained from the process b. The starting 80 materials used for process b are in some cases known, but in most cases unknown. These unknown starting materials may, however, be obtained according to processes known per se.

Starting materials of the formula II

$$R^{2}$$

$$R^{2}$$

$$R^{3}$$

$$R^{4}$$

$$R^{5}$$

$$R^{5}$$

$$R^{5}$$

wherein Z<sup>1</sup> is SH may be obtained from the corresponding o - phenylenediamine by reaction with potassium ethylxanthate (Org. Synth. Vol. 30, p. 56) or thiophosgene.

The compounds of the formula II wherein Z<sup>1</sup> is alkylmercapto and alkylsulfinyl may be obtained from the above mentioned compound by simple Salkylation with alkyl halide and by oxidation of the product from the S - alkylation, respectively.

95

The compounds of the formula II wherein Z1 is halogen or acyloxy may be obtained from compounds of the same formula wherein Z1 is OH by treatment with POCl<sub>3</sub>, POBr<sub>3</sub> and the like or the appropriate acyl halide, respectively. The starting 100 material wherein Z1 is OH is obtained from the corresponding o - phenylenediamine by reaction with phosgene.

The o-phenylenediamines required may be obtained from the corresponding substituted ben-105 zenes according to processes known per se, e.g. by the consecutive processes: nitration, reduction, acetylation, nitration, deacetylation and reduction, or from one of the intermediary stages just mentioned. In order to obtain a o - phenylenediamine wherein R<sup>5</sup> 110 is other than H, acylation (by the group R14CO) is preferably made on the nitro - aniline stage.

Starting materials of the formula

wherein R15 is H, may be obtained either from the correspondingly substituted (R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup>) 2 - methyl - substituted pyridine N - oxide via a known rearrangement to the intermediate 2 - pyridinylmethanol 5 urvia a hydroxymethylation of the substituted (R<sup>6</sup>, R<sup>7</sup> and R8) pyridine to give the same intermediate, and then treatment of the 2 - pyridinylmethanol with halogenating agents such as thionyl chloride or Oacylating agents such as p - toluenesulfonyl chloride 10 to give compounds of the formula III wherein  $Z^2$  is halogen and sulfonyloxy groups, respectively.

These leaving groups may then be substituted for alkylmercapto groups by treatment with e.g. sodium alkylmercaptide, which may then be oxidized to an 15 alkylsulfinyl group, or substituted for SH by treatment with e.g. NaSH.

For the preparation of intermediates of formula

wherein R7 is alkoxy, alkenyloxy, alkynyloxy, alkoxyalkoxy and dialkylaminoalkoxy, a compound of 20 formula VII, wherein R7 is NO2, is reacted by the corresponding sodium alkoxide. Analogously, for the preparation of an intermediate of formula VII wherein R<sup>6</sup> and R<sup>7</sup> or R<sup>7</sup> and R<sup>8</sup> form a ring structure including an oxygen atom at position 4, a compound of formula 25 VII wherein R7 is NO2 and R6 or R8 represents

hydroxyalkyl is reacted with a non-nucleophilic base. The following intermediates A) and B) are included in the scope of the invention:

A) New compounds of the formula

30 wherein  $R^{1a}$ ,  $R^{2a}$ ,  $R^{3a}$  and  $R^{4a}$  are the same or different and selected from the groups

(a) H,

(b) alkyl containing 1-6 carbon atoms, including cycloalkyl,

(c) alkoxyalkyl containing 1-3 carbon atoms in the alkoxy part and 1-6 carbon atoms in the alkyl part,

(d) aryloxyalkyl containing 1-3 carbon atoms in the alkyl part.

(e) arylalkyl containing 1-6 carbon atoms in the 40 alkyl part,

(f) aryl,

(g) alkoxy containing 1-6 carbon atoms,

(h) alkoxyalkoxy containing 1-3 carbon atoms in the outer part and 1-6 carbon atoms in the part 45 nearest the aromatic ring,

(i) aryloxyalkoxy containing 1-6 carbon atoms in the alkoxy part,

(j) arylalkoxy containing 1-6 carbon atoms in the alkoxy part and

(k) aryloxy,

R<sup>50</sup> is

(a) H,

(b) alkoxycarbonyl containing 1-4 carbon atoms in the alkoxy part,

(c) arylalkoxycarbonyl containing 1-2 carbon atoms in the alkoxy part,

(d) dialkylaminocarbonyl containing 1-4 carbon atoms in each alkyl group, or

(e) arylamine: irbonyl,

and  $Z^{1a}$  is

(a) SH.

(b) Clor Br

and provided that not more than one of R10, R20, R30 and  $R^{4a}$  is H, are suitable intermediates for the 65 preparation of compounds of the formula I with R1,  $R^2$ ,  $R^3$ ,  $R^4$  and  $R^5$  having the same meaning as  $R^{1a}$ ,  $R^{2a}$ ,  $R^{3a}$ ,  $R^{4a}$  and  $R^{5a}$ , respectively, according to method b.

B) New compounds of the formula

wherein R<sup>60</sup> and R<sup>80</sup> are

(a) Hor

(b) alkyl containing 1-5 carbon atoms, and R7a is

(a) alkenyloxy containing 2-5 carbon atoms, or

(b) alkynyloxy containing 2-5 carbon atoms,

(c) oxacycloalkyl containing one oxygen atom and

75 3-7 carbon atoms

(d) oxacycloalkoxycontaining two oxygen atoms and 4-7 carbon atoms

(e) oxacycloalkylalkyl containing one oxygen atom and 4-7 carbon atoms

(f) oxacycloaikylalkoxy containing two oxygen atoms and 4-6 carbon atoms,

(g)  $R^{6a}$  and  $R^{7a}$ , or  $R^{7a}$  and  $R^{8a}$  together with the adjacent carbon atoms in the pyridine ring form a ring wherein the part constituted by  $R^{6a}$  and  $R^{7a}$  or  $R^{7a}$  and

-O-(CH<sub>2</sub>)<sub>pa</sub>-

—CH<sub>Z</sub>—(CH<sub>1pa</sub>—

-O-CH=CH-

wherein pais 2, 3 or 4 and the O atom always is attached to position R7a, and Z2 is

(a) SH,

(b) halogen CI, Br, I or

and provided that not more than one of R<sup>6a</sup> and R<sup>8a</sup> is H, are suitable intermediates for the preparation of compounds of the formula I with R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> having 100 the same meaning as R<sup>6a</sup>, R<sup>7a</sup> and R<sup>8a</sup>, respectively,

according to method b.

For clinical use the compounds of the invention are formulated into pharmaceutical formulations for oral. rectal, parenteral or other mode of administration.

unreacted starting material. The oil was chromatographed on a silica column using CH<sub>3</sub>OH—CH<sub>2</sub>Cl<sub>2</sub> 5:95 as eluant and then the product was recrystallized from CH<sub>3</sub>CN giving the desired product in crystalline 5 form (0.85 g, 32%), m.p. 116°C.

Which one of these two procedures that have been used for the preparation of the different sulfoxides have been indicated in Table 2 below. For most of the compounds synthesized according to example 2 the 10 chromatographic separation was not performed.

Example 3. Method b. Preparation of 4,6 - dimethyl-5-methoxy-2-[[(3,4 - dimethyl-2 - pyridinyl) methyl] thio] - 1H - benzimidazole.

To 4,6 - dimethyl - 5 - methoxy - 2 - mercapto - 1H15 benzimidazole (1.04 g, 0.0050 mol) in methanol (50 ml) were added (in the following order) NaOH (0.2 g, ;.0050 mol) dissolved in water (2 ml) and 3,4 - dimethyl - 2 - chloromethylpyridine hydrochloride (0.96 g, 0.0050 mol). The mixture was heated until reflux. NaOH (0.2 g, 0.0050 mol) dissolved in water (2 ml) was added dropwise and then the reflux was continued for 3 hours. The mixture was poured on ice-water (200 ml). Filtration and recrystallization from CH<sub>3</sub>CN gave the desired product (1.1 g, 67%).

25 NMR data for the final product is given below.

Example 4 and 5. Method d. Preparation of N¹benzoyl - 5 - methoxy - 2 - [[(4 - methoxy - 3,5 dimethyl - 2 - pyridinyl) methyl] - thio] - 1Hbenzimidazole and N¹ - benzoyl - 6 - methoxy - 2 - [[(430 methoxy - 3,5 - dimethyl - 2 - pyridinyl) methyl] thio] 1H-benzimidazole

1H-benzimidazole
5-Methoxy-2-[[(4-methoxy-3,5-dimethyl-2-pyridinyl) methyl]-thio]-1H-benzimidazole (3.0 g, 0.009 mol) was dissolved in CH<sub>3</sub>CN (30 ml) and
35 triethylamine (1.9 ml) was added. Benzoyl chloride (1.4 g, 0.010 mol) was added dropwise under stirring during 15 min. Then the mixture was stirred at 55°C for 45 min. The solvent was evaporated off and ether was added to the residue under ice-cooling. The crystalline residue, thus obtained was stirred with water, filtered off and dried giving a white crystalline product mixture (1.9 g, 48%) of the desired two products in a 75:25 molar ratio (according to HPLC-analysis and NMR). NMR data for the final products is

Example 6. Method d. Preparation of N - methoxy-carbonyl - 5.6 - methylenedioxy - 2 - [[(4 - methoxy - 3.5 - dimethyl - 2 - pyridinyl) - methyl] sulfinyl] - 1H-benzimidazole.

50 Chloro methylformate (0.24 g, 0.0026 mol) dissolved in CH<sub>2</sub>Cl<sub>2</sub> (5 ml) was added dropwise to a stirred solution of 5,6 - methylenedioxy - 2 - {[(4 - methoxy - 3,5 - dimethyl - 2 - pyridinyl) - methyl] sulfinyl] - 1H- benzimidazole (0.80 g, 0.0022 mol) and triethylamine in CH<sub>2</sub>Cl<sub>2</sub> (10 ml). The mixture was then stirred at room temperature for 19 h. The CH<sub>2</sub>Cl<sub>2</sub>-solution was washed with water, dried (MgSO<sub>4</sub>) and the solvent was evaporated giving the desired product as an oil (0.06 g, 6%). NMR data for the final product is given below.

Example 7. Method d. Preparation of  $N^1$  - (N' - phenylcarbamoy!) - 5,6 - methylenedioxy - 2 - [[(4 - methoxy - 3,5 - dimethyl - 2 - pyridinyl) - methyl] sulfinyl] - 1H- benzimidazole.

Phenylisocyanate (0.20 g, 0.00167 mol) dissolved in CH<sub>2</sub>Cl<sub>2</sub> (5 ml) was added dropwise under stirring to a solution of 5,6 - methylenedioxy - 2 - [[(4 - methoxy - 3,5 - dimethyl - 2 - pyridinyl) - methyl] sulfinyl] - 1H - benzimidazole (0.50 g, 0.00139 mol) and triethylamine (0.28 g, 0.00278 mol) in CH<sub>2</sub>CL<sub>2</sub> (15 ml). The mixture was then stirred at room temperature for 50 hours. The CH<sub>2</sub>Cl<sub>2</sub>-solution was washed with water, dried (MgSO<sub>4</sub>) and the solvent was evaporated giving the desired product as an oil (0.03 g, 5%). NMR data for the final products is given below.

Example 8. Method e. Preparation of 4,6-dimethyl - 5-methoxy - 2 - [[(4 - methoxy - 3,5 - dimethyl - 2 - pyridinyl]methyl]sulfinyl] - 1H - benzimidazole.

N¹ - Propionyl - 4,6 - dimethyl - 5 - methoxy - 2 - [[(4 - methoxy - 3,5 - dimethyl - 2 - pyridinyl)methyl]sulfinyl] - 1H - benzimidazole (1.0 g, 0.0023 mol) was heated in 1M NaOH (15 ml) for 1 h under stirring and N<sub>2</sub>-atmosphere, pH was adjusted to 9.5 by addition of 2M HCI. Extraction with CH<sub>2</sub>Cl<sub>2</sub>, separation of the

85 phases, drying the organic phase, evaporation of the solvent and recrystallization from CH₃CN gave the desired product (0.30 g, 35%), m.p. 137°C.

The following Table 2 gives data for further examples of compounds of the invention.

Table 2. Summary of working examples.

9 <sup>7</sup>	, i
R R 15	N \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \
<sup>2</sup> γ cu-x	《人人》。
	15 Y <sub>4</sub> "

L A	ı	a <sup>15</sup>	R <sup>1</sup>	R <sup>2</sup>		R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	g <sup>6</sup>	R <sup>7</sup>	a <sup>S</sup>	Method {[x, No.}	Yield 1	M.p.( <sup>D</sup> C) other data
•	<u> </u>	н	Cm3	Ch.	<u> </u>	CHY	CH <sub>1</sub>	н	CH3	осн <sub>2</sub> сн•сн <sub>2</sub>	СНЗ	b (Ex 3)	82	164-165
			CHY			CH <sub>2</sub>	ᅄ	H	CH3	OCH2CH-CH2	сн3	a (Ex 2)	73	1/6-148
			CH	Ct.,		CHZ	CH <sub>3</sub>	н	CH <sub>3</sub>	осн <sub>3</sub>	CH3	p (E= 3)	79	207
	50		CH <sub>1</sub>	CH <sub>2</sub>		•	CH <sub>3</sub>		CH3	осн	CH3	a (E = 2)	32	193
			CH	•		•	,		OH <sub>3</sub>	OCH <sup>2</sup> CH+CH <sup>2</sup>	CH3	5 (E = 1)	97	165
	50		CH3	-		ر پېن	н	н	CH3	OCH2CH+CH2	CH3	* (E # 2)	59	147
			CH			Cri,		н	CH,	осн <sub>1</sub>	CH3	5 (E + 3)	79	159
	50		CH <sub>1</sub>	•		CH,		н	CH3	осн <sub>3</sub>	CH3	a (Ex 1)	83	188
			CHI	•	ı	•	CH3	н	CH3	OCH <sup>2</sup> CH+CH <sup>2</sup>	CH3	b (Ex 3)	77	NMR

(:	1	215	g)	n²	ر۽	<b>n</b> <sup>4</sup>	<b>8</b> 5	**	47	R <sup>5</sup>	Method {[s. No.]	field 1	H.p.( <sup>0</sup> () other data
18	54	H	٥'n	013	М	CH3	н	CH3	001201-012	CH <sub>3</sub>	a (E= 1)	54	129
19	s	M	OL <sub>3</sub>	OI <sub>3</sub>	>	CH3	Ħ	OI3	0013	CH <sub>3</sub>	b (Ex 3)	79	163
20	so	ж	Oi <sub>3</sub>	CH <sub>3</sub>	н	O13	н	CH <sub>3</sub>	осн <sub>3</sub>	CH3	a (Ex 1)	52	191
21	s	M	Or <sub>3</sub>	013	н	н :	Ħ	CH <sub>3</sub>	OCH <sub>2</sub> CH+CH <sub>2</sub>	CH3	b (Ex 3)	37	109
22	50	н	Oi <sub>3</sub>	OI,	н	H	н	O13	001201-012	CH3	a (Ez 1)	58	149
23	5	н	×	CH	CH <sub>3</sub>	H	н	CH <sub>3</sub>	0CH2CH+CH2	CH3	b (Ex 3)	99	181
24	50	н	M	CN <sub>3</sub>	OI,		н	CH <sub>3</sub>	001201+012	CH3	a (Ex 1)	71	157
25	s	M	CH <sub>2</sub>	×	×	CH <sub>3</sub>	н	CH <sub>3</sub>	001201-012	013	b (Ex 3)	62	1017
26	so	N	CH <sub>2</sub>	н	×	CH3	н	DI <sub>3</sub>	001201-012	CH3	a (Ex 1)	10	155
27	5	H	OI3	Ħ	H	н	н	CH <sub>3</sub>	001201-012	CH3	b (Ex 3)	90	104R
28	so	16	013	н	ж	н	н	CH <sub>3</sub>	OCH2CH+CH2	013	a (Ex 1)	69	142
25	s	н	×	CH <sub>3</sub>	H	×	н	CH <sub>3</sub>	0CH2CH-CH2	CH <sub>3</sub>	b (Ex 3)	74	MAR
30	50	н	м	Di <sub>λ</sub> .	н	н	н	CH <sub>3</sub>	OCH_CH+CH2	043	a (Ex 1)	55	134
31	5	н	10	0013	×	H	н	CH3	OCH_CH-CH <sub>2</sub>	OI3	b (Ex 3)	51	105-107
32	so	м	н	OCH <sub>3</sub>	н	н	H	CH <sub>3</sub>	OCHZON-CHZ	CH <sup>3</sup>	a (Ex 1)	62	111
33	s	Ħ	н	оси <sub>3</sub>	н	H	H	O13	OCH <sup>2</sup> CE CH	CH3	b (Ex 3)	64	154
											<del></del>		cont

cont.

E1	x	R <sup>15</sup>	R <sup>1</sup>	R <sup>2</sup>	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>	Method ({z. No.}	Yield 1	M.p.( <sup>O</sup> C) other data
34	so	rt	×	OCH,	н	н	н	CH3	OCH <sub>2</sub> CICH	··13	• (Ex 1)	71	145
35	so		H	0CH <sub>2</sub>	н	н	H	<b>:</b> 1	OCH 3	c <sub>z</sub> u <sub>s</sub>	a (Ex 1)	31 -	147
36	s	н .	н	0СН3	н	H	н	×		-(CH <sub>2</sub> )4-	b (Ex 3)	61	ME
37	so		н	осн3	~×	н	H	н		-(CH <sub>2</sub> ) <sub>4</sub> -	• (Ex 2)	ж	MER
38	s	н	н	$(\mathcal{C}_{0}^{\circ})$	. н	н	Ħ	CH <sup>3</sup>	осн3	CH3	b (Ex 3)	22	148
40	s	H	OI <sub>3</sub>	н	CH3	н	н	CH3	OCH2CH+CH2	CH <sub>3</sub>	b (Ex 3)	76	134-136
41.	so	н	CK1	h	СН,	н	H	ᅄ	0CH2CH=CH2	CH3	a (Ex 1)	35	111
42	s	н	н	OCH,CN	н	H	H	CH3	осн <sub>3</sub>	CH3	b (Ex 3)	29	66
43	so	н	н	осн,сн	н	н	н	CH <sub>3</sub>	0CH <sub>3</sub>	CH3	* (Ex 1)	39	- 94
44	s	н	×	$\langle \rangle$	н	н	н	CH3	осн <sup>3</sup>	, c <sup>ici</sup> a	b (Ex 3)	75	HMR
45	SO	н	×	$\bar{\bigcirc}$	н	H	н	CH3	OCH3	CH3	a (Ex 2)	60	155

Ĺ	,3	R <sup>15</sup>	<b>R</b> <sup>1</sup>	# <sup>2</sup>	g <sup>3</sup>	g <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>S</sup>	Method · (Ex. No.)	Tield Z	M.p.(°C) other data
47	50		×	(0001,01,001,	CH,	н	H	CH3	осн <sub>3</sub>	CH3	•		
48	s	Ħ	M	cochi O	O+3	×	H	CH <sub>3</sub>	осн <sub>3</sub>	CH3	¢		
49	50	H	M	COOCH TO	ᅄ	H	H	CH;	∞н3	CH3	•		
50	s	н	*	CrizON	ᅄ	H	н	CH3	осн3	CH3	b (Ex 3)	86	192
51	<b>SO</b>	Ħ	п	CHZON	O13	н	H	OI3	оси 3	CH <sub>3</sub>	a (Ex 1)	10	169
52	\$	M	u 7	CH <sup>2</sup> OCO-	ᅄ	H	×	CH3	OCH <sup>3</sup>	CH3	¢		
53	50	×	M	CH <sub>Z</sub> 0CO-(())	CH.	н	H	CH <sub>3</sub>	осиз	СНЗ	•		
54	S	H	h	COOCH <sup>3</sup>	· он <sub>з</sub>	H	н	CH3	0CH <sup>2</sup> CH-CH <sup>2</sup>	CH3	b (Ex 3)	75	168
55	SO	H	×	соосн	CH3	H	н.	CH3	OCH <sup>2</sup> CH-CH <sup>2</sup>	CH3	a (Ex 1)	52	139
56	\$	×	Cr3	0CH <sub>3</sub>	CN 3	H	н	CH3	0CH <sub>3</sub>	CH3	b (Ex 3)	70	HER
8	SO	Ħ	CH <sub>3</sub>	осн <sub>3</sub>	CH3	H	H	CH3	OCH <sub>3</sub>	CH <sub>3</sub>	(a (E= 1)	35	133
3	s	Ħ	013	осж <sub>3</sub>	O13	Ħ	н	CH3	OI3	н	b (Ex 3)	67	INT
1	so	m	CH <sub>3</sub>	0013	O13	×	H	CH3	CH <sub>3</sub>	н	a (Ex 1)	32	161
57	5	×	O13	оси <sub>г</sub> си <sub>г</sub> оси <sub>з</sub>	CH <sub>3</sub>	н	н	CH <sub>3</sub>	осн <sub>3</sub>	CH3	b (Ex 3)	90	200
58	50	×	CH <sub>3</sub>	OCH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>	Сн3	H	H	CH3	осн <sub>3</sub>	CH3	a (Ex 1)	-68	144

ĹZ	x	R <sup>15</sup>	a <sup>1</sup>	A <sup>2</sup>	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>B</sup>	Method (Ex. No.)	Tield 1	M.p.( <sup>O</sup> C) other data
59	s	н	CH3	OCH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>	CH <sub>3</sub>	н	н	н	СНЗ	CH <sub>3</sub>	b (Ex 3)	95	IMR
60	sc	H	O13	00H20H20CH3	ОН,	н	н	ж	CH <sup>3</sup>	. CH <sub>3</sub>	a (Ex 1)	58	131
61	5	и	CH <sub>3</sub>	COCH3	CH <sub>3</sub>	н	н	CH3	осн <sub>3</sub>	CH <sub>3</sub>	b (Ex 3)	90	192-4
62	sc	) н	CH <sub>3</sub>	COCH <sub>3</sub>	CH <sub>3</sub>	н	::	CH3	OCH3	CH3	a (Ex 2)	25	164-5
63	·s	н	uiz	LOCH <sub>3</sub>	CH3	н	H	CH3	H.	CH3	b (Ex 3)	99	184-6
64	sc	) ×	CH3	сосн	CH <sub>3</sub>	н	н	CH3	н	CH <sub>3</sub>	a (La. 2)	ຍ	148-50
65	:	N	CH <sub>3</sub>	COCZHS	CH3	н	Ħ	CH3	осн 3	CH3	b (Ex 3)	68	. 149
64	SC	Эн	CH <sub>3</sub>	COCZNS	CH <sub>3</sub>	н	н	CH3	осн <sub>3</sub>	СНЗ	a (Ez 2)	48	INTR .
67	s	н	CH <sub>3</sub>	C2H5	O13	н	н	CH3	OCH <sub>3</sub>	CH <sub>3</sub>	b (Ex 3)	91	182
68	SC	3 н	CH3	C <sub>2</sub> H <sub>S</sub>	CH3	H	н	CH3	осн	CH <sub>3</sub>	a (Ex 2)	67	175-7
69	s	н	CH <sub>3</sub>	Carlo	CH <sub>3</sub>	н	н	CH3	осн <sub>3</sub>	H	b (Ex 3)	95	10·R
70	SC	э ж	CH <sub>3</sub>	CzHs	О13	н	н	CH3	осн 3	H	• (Ex 2)	73	142-3
71	s	н	C <sub>2</sub> H <sub>5</sub>	ĊN	C <sub>2</sub> H <sub>2</sub>	н	н	сн,	осн <sub>3</sub>	CH3	b (Ex 3)	82	150
72	S	эн	CzHs	CH	C <sub>Z</sub> H <sub>S</sub>	н	н	CH3	осн <sub>3</sub>	Сн3	a (Ex 2)	81	180
73	s	н	CH	OCH <sub>3</sub>	СН,	И3	н	CH <sub>3</sub>	осн <sub>3</sub>	CH3	b (Ex 3)	82	143
74	SA	ОН	, CH <sub>3</sub>	OCH 3	CH <sub>3</sub>	ᅄ	H	CH <sub>3</sub>	0СН3	CH3	a (Ex 2)	43	163 -

L A		415	41	4 <sup>2</sup>	43	g4	R <sup>5</sup>	<b>A</b> 6	R <sup>7</sup>		Method ((s. No.)	Tield 1	M.p.( <sup>0</sup> () other data
75	<u> </u>	,	C1	CI	CI	н	м.	CH <sub>3</sub>	OCH <sub>3</sub>	CH3	b (C= 3)	90	204
	50		C1	C1	CI	н	н .	CH <sub>3</sub>	OCH <sub>3</sub>	CH3	•		
77		n	н	CH <sub>3</sub>	CH3	н	н	н ,	осн <sub>3</sub>	C <sub>2</sub> H <sub>5</sub>	a (E = 1)	. 43	156
78	s	H	н	cos C	н	н	H	CH3	осн3	ᅄ	b (Ex 3)	90	IPR
79	50	н	н	CM2	H	н	. #	CH)	0CH3	CH3	a (Ex 1)	61	MR
80	s	м	м	-осн <sub>7</sub> 0-		×	16	CH <sub>3</sub>	0013	CH3	b (Ex 3)	91	168
81	•		и	-оси <sub>2</sub> о-		н	ж	СН,	OCH,	CH <sub>3</sub>	a (Ex 1)	67	165
82	5	×		:H-CH+CH-	н	н	н	•	осн <sub>3</sub>	CH <sub>3</sub>	b (Ex 3)	73	MAR
e 83	•	 H		CH-CH-CH-		н	н	CH <sub>2</sub>		CH <sub>2</sub>	a (Ex 1)	60	. 184
			H	-OI-CI-CI-CI	. "	н	H	CH3/	•	CH <sub>1</sub>	b (Ex 3)	78	191
84		4		-01-01-01-01-		н	H	CH	0CH <sub>3</sub>	CH <sub>3</sub>	a (Ex 1)	34	175
	SO.		H		н	ж		CH	оси <sub>3</sub>	CH	b (Ex 3)	58	10·00
86		н	•	CH2CH2CH2-	н	. н	 **	ci,	0CH <sub>2</sub>	ભં	a (Ex 1)	27	175
87		Ħ	•	CH2CH2CH2-	п			4.3	0013	CH <sub>3</sub>	d		
88	s	H	н	-00120-			CO <sub>2</sub> CH <sub>3</sub>	CH <sub>3</sub>	- · · · · · · · · · · · · · · · · · · ·				cont

 [ a	1	R <sup>15</sup>	R1	· <sub>R</sub> 2	R <sup>3</sup>	R <sup>4</sup>	R.		R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>	Me thod (Ex. No.)	Tield 2	M.p.( <sup>O</sup> C) other dat
6	so	и	H	-0CH <sub>Z</sub> 0-		н	C	2013	머	оси <sub>3</sub>	CH3	d (Ex 6)	6	MA
7	so	н	н	-осн <sub>2</sub> 0-		H	COM	<b>.</b> @	ᅄ	осн	CH3	d (Ex 7)	5	104R
90	5	H	H	0CH,CH2CH20-	н	H	. н		CH3	осн3	CH3	b (Ex 3)	25	NMR.,
91	50	н	n	OCH2CH2CH2O	н	н	н	;	CH3	осн <sub>3</sub>	CH <sup>3</sup>	a (Ex 2)	78	61
92	s	н	CHZ	0(CH <sub>2</sub> ) <sub>6</sub> CH <sub>3</sub>	OI,	н	H		CH <sub>3</sub>	осн <sub>3</sub>	O13	b (Ex 3)	64 ·	NIR
	50		CH	0(CH <sub>2</sub> )6CH <sub>3</sub>	ᅄ		H		CH3	OCH <sub>1</sub>	CH <sub>3</sub>	a (Ex 2)	32	116
	s		ч.	C <sub>2</sub> H <sub>5</sub>	H	н	H		CH3	OCH <sub>2</sub> CH=CH <sub>2</sub>	CH3	b (Ex 3)	45	XMR
	so		н	د ع د <sub>ک</sub> بار	н	н	н		CH3	OCH <sub>2</sub> CH-CH <sub>2</sub>	CH <sub>3</sub>	a (Ex 1)	49	124-6
95		н	н	0CH <sub>3</sub>	×	н			CH <sub>3</sub>	0CH2CH2CH(CH3)2	CH <sub>3</sub>	b (Ex 3)	\$\$	MAR
96		н		осн <sub>3</sub>	н	н	H	l	CH3	OCH2CH2CH(CH3)2	CH <sub>3</sub>	# (Ex 1)	33	111
97		н				н	,	ı	CH3	0CH <sup>3</sup>	CH3	b (Ex 3)	96	190
98	-	H				н	H	,	. сн <sub>3</sub>	0CH <sub>3</sub>	CH3	a (Ex 2)	93	109
ė	5	H	н	осн,	н	н	CC	<del>(</del> 0)	) CH <sub>3</sub>	осн3	CH3	d (E= 4)}	48	1942
5	s	н	н	н	ОСН	, H	CC	-√Ö	СН3	осн <sub>3</sub>	ᅄ	d (Ex 5)∫		

Table 2 comt.

Ĺs.	I	R <sup>15</sup>	<b>A</b> 1	a <sup>2</sup>	# <sub>3</sub>	84	R <sup>S</sup>	<b>x</b> <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>	Method (Ex. No.)	Yield S	H.p.( <sup>O</sup> C) other data
99	5	И	н	O((O(3)2	И	H	н	CH3	001201-012	CHJ	b (Ex 3)	99	70
101	\$	M	ĸ	c(O(3)3	И	М	H	CH3	001201-01 <sup>2</sup>	CH <sub>3</sub>	b (Ex 3)	52	88-89
102	50	Ħ	H	C(CP1 <sub>2</sub> ) <sub>3</sub>	H	M	×	CH3	001201-012	сн <sub>э</sub>	4 (Cx 2)	12	IMA
103	S	H	M	נוסקיסלוס	И	ĸ	н	СЧ	0CH <sub>3</sub>	ᅄ	b (E= 3)	84	NAC .
104	<b>50</b>	H	M	נוספלומלום	. *	M	<b>H</b> .	CH3	0013	CHO	a (Es 1)	38	118
105	S	M	×	\$	Ļ	×	×	CH3	0CH <sub>3</sub>	CH3	b (Ex 3)	58	216
105	so		ĸ	ζ	$\mathcal{L}$	Ħ	n .	CH3	осн <sub>3 .</sub>	CH3	a (Ex 2)	32	158
107	50	H	H	0CH3 -0	W	M	رات <sub>ك</sub> ش	CH3	ocu3	снэ	d) (Ex 4 and 5	) {e	{ Nest
106	59	Ħ	M	` <b>x</b>	<b>60</b> 1	н.	co <sub>z</sub> ci <sub>j</sub>	CH3	oci <sub>l</sub>	ᅄ	[ه	`{	l
109	5		H	soi <sub>3</sub>	10	×	M	CH <sub>3</sub>	0CH <sup>3</sup> .	. <sup>O1</sup> 3	b (Ex 1)	83	147-148
110	S	W	M	OI(OI3)2	×	H	×	013	OCH 2-	CH3	b (Ca 3)	86	JN MAK
111	<b>24</b>	*	h	си(си <sup>3</sup> ) <sup>3</sup> .	×	H .		СНЗ	OCH E	CH <sub>3</sub>	a (Ex 2)	89	<sup>1</sup> H MMR

cont.

Table 2 cont

(a I	215	RI	R <sup>2</sup>	83	R <sup>4</sup>	A2	R <sup>5</sup>	R <sup>7</sup>	RB	Method (Ex: No.)	Tield Z	M.p.(°C) other data
112 5	н	и.	CIL,CIL,COOI,	н	N	н	CH3	OCH <sup>2</sup> CH-CH <sup>2</sup>	сн3	b (Ex 3)	40	. <sup>1</sup> H JUR
113 50	×	H	cijaljcoci	н	H	H	CH3	OCH <sup>2</sup> CH+CH <sup>2</sup>	CH3	a (E= 2)	28	123-4
114 \$	×	H	دره ً	×	N	н	CH3	осн3	CH3	b (E= 3)	21	162
115 5	И	H	001	×	H	H	-CH+	CH-CH-CH-	Ħ	b (Ex 3)	67	105
116 50	H	M	осн <sub>3</sub>	M	Ħ	×	-CH-	си-си•си-		4 (Ex 1)	66	100
117 S	н	н	•-⊘	×	H	н	СНЗ	оснэ	см3	b (Ex 3)	98	122
118 SO	н	×	<b>∘</b> -⊘	×	ж	н	Сн3	осн <sub>3</sub>	CH3	a (2c2)	80	118
119 \$	Ħ	11	0CH2CH2	×	M	H	CH.3	осн3	CH3	b (Ex 3)	80	1 <sub>H</sub> 1048
120 50	×	H	∞ئەتئەن	×	H	H	CH3	OCH <sup>3</sup>	CH3	* (Ex 5)	55	145 d
121 S		×	<b>∞</b> - <b>⊘</b>	×	M	M	CH3	осн3	c <sub>H</sub> 3	6 (E= 3)	82	1H MAR
122 50	M		<b>u</b> -⊘	н	N	H	СнЭ	осн <sub>3</sub>	Сн3	a (Ex 2)	24	1H HHR
123 S	ĸ	m	<b>-</b> ⊘	×		n	CH3	осн3	CH3	b (Ex 3)	88	158

cont.

En x n	5 ,1	R <sup>2</sup>	• 3	n*	R 5	10	a <sup>7</sup>	k <sup>S</sup>	Method [s. No.)	field 3	M.p. ('C) other data
124 SO N	n	-⊘	H	×	<b>*</b>	СНЗ	œ <sub>*</sub> j	сн3	a (Ex 2)	52	104
125 S 18	n	SOCH <sub>3</sub>	H	н	н	CH3	осн <sub>3</sub> .	СНЗ	b (Ex 3)	57	ln mar
126 SO H	н	SOCH <sub>3</sub>	*	н	н .	CH)	осн <sub>3</sub>	CHO	a (Ex 1)	47	, I India
127 SO N	н	NO,	×	н	н	CH3		C#3	a (Ex 1)	14	N MAG
128 S H	H	Br	×	H	×	Сн	оси <sub>2</sub> си-си <sub>2</sub>	CH3	b (2x 3)	64	171
129 SO H	H	Br	и	н	н	CH <sub>3</sub>	оси2си-си3	CH3	a (Ex 2)	56	143
130 S H	н	осн <sub>3</sub>	H	H	H	-CH+C	H-0-	н	b (Ex 3)	77	1948
131 SO H	н	оси3	H	н	н	-CH-C	н-0-	H	a (Ex 2)	19 -	
132 SO N	H	СНЭ	си <sub>э</sub>	H	SOC (CH <sup>2</sup> )	3 CH3	осн <sub>3</sub>	CH3	d (Ex 6)	22	T68
134 SO H	×	CH <sub>3</sub>	. сн3	н	Sucoi3)5	СНЗ	осн	CH <sub>3</sub>	d (Ex 6)	21	<sup>1</sup> )+ 1046
135 S H	M	CH3	CH3	H	н .	Сн	OCH 2 O	CH3			
136 SO H	N	CH3	CHJ	H	н	Сн	OCH-	CH3		•	
137 S H	н		-EH2CH2C42-	н	H	LH3	осн	CH3	b (Ex 3)	74	160
138 SO H	н		-CH <sub>2</sub> CH <sub>2</sub> CH <sub>2</sub> -	н	н	CH3	оси	CH3	a (Ex 1)	40	171

Table 2 cost

Ex 1	ı	2 <sup>15</sup>	R <sup>1</sup>	R <sup>2</sup>	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>	Method {Ex. No.}	Tield Z	M.p. (°C) other data
139 5		38	-CH-	CH-CH-M-	Я	И	н	. СИ3	оси3	CH3	b (Ex 3)	- 38	1948 -
140 5	so	H	-CH-	CH-CH-N-	н	н	н	CH3	осн	CH3	• (E= 1)	26	60
141 :	5	и	н		-0CH <sub>2</sub> 0-	н	H	CH <sub>3</sub>	СНЗ	CH3	b (E= 3)	. 83	193-95
142 9	50	×	H	a	OCH <sub>2</sub> 0	H	H	CH3	сн3	CH3	a (Ex 2)	76	173
143 5	50	12	H	COCH	CH <sub>3</sub>	×	ж	H	OCH3	C2H5	a (Ex 2)	49	154
144 5		ж	CH <sub>3</sub>	•	CH <sub>3</sub>		H	СНЗ	CH <sub>2</sub>	н	b (Ex 3)	39	THE HAR .
145 5			CH3	•	OI,		н	CHZ	CH3	, ж	a (Ex 2)	65	TH JUST
146 5		*	CH.	-	CH <sub>3</sub>		н	и	CH3	CH3	b (Ex 3)	78	143
147 5			CH	•	CH <sub>1</sub>		н	- н	CH3	CH3	a (Ex 2)	64	180
148 5			CH <sub>3</sub>	-	СН <sub>3</sub>		H	СНЗ	H	CH <sub>3</sub>	b {Ex 3}	70	239-42
49 5			CH <sub>2</sub>	•	CH <sub>3</sub>		н	СН.	H	CH <sub>3</sub>	a (Ex 2)	14	171
150 5			CH <sub>1</sub>	•	н	СНЭ	н	CHI	CH3	н	b (Ex 3)	96	210
150 S			-	•	н	CH3		CH3	CH <sub>3</sub>	н	a (Ex 2)	66	<sup>3</sup> н ючя
			CH <sub>2</sub>	•	C#3	•	н	•	OC <sub>2</sub> H <sub>5</sub>	СНЗ	b (Ex 3)	94	151
			-		•3 СН <sub>3</sub>		н	•	· 0С <sub>2</sub> н <sub>5</sub>	cii	1 (Ex 2)	29	150
154 :		л н	СH <sub>3</sub>	<u></u>	7′ H	н	н	н .	сн <sub>3</sub>	C <sub>2</sub> H <sub>5</sub>	b (Ex 3)	48	1 <sub>H KN-R</sub>

cont.

Table 2 cont.

(a z a	5 g1	R <sup>2</sup>	A <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup> R <sup>7</sup>	R <sup>8</sup>	Mehtud (Ex. No.)	Tield	M.p. ('C) other data
155 SO H	и	$\overline{\langle}$	ÿ н	И	н	н сн	C2H5	a (Ex 2)	44	105
15 <b>6</b> S H	н	$-\overleftarrow{\bigcirc}$	*	H	н	снз оснаснаосн	СН3	b ([= 3)	94	l mar
157 SO H	×	$\prec >$	H	H.	н	CH3 OCH2CH2OCH	сн3	a (Ex 2)	18	181
158 S H	н	σ,	н	M	н	CH3 OCH2 0	СН3	b (E x 3)	67	100
159 SO N	м	CF <sub>3</sub>	н	H	. н	сн3 осн5	CH3	a (Ex 2)	57	125
160 S N	H	CHZCHZCOOCZHS	• н	н	H	снэ оснэ	CHO	b (Ex 3)	15	IH WIR
161 SO H	×	OCH <sub>3</sub>	H	н	č-0010	н <sub>3</sub> ) <sub>3</sub> сн <sub>3</sub> осн <sub>3</sub>	CH3	d (Ex 6)	50	155
163 SO N	н	OCH <sub>2</sub>	М	M	н	-сн <sub>2</sub> сн <sub>2</sub> 0-	H			
164 S 44	н	OCH ,	. 11	H	н	-си <sub>2</sub> си <sub>2</sub> си <sub>2</sub> 0-	н	b (Ex 3)	71	H NOOR
165 SO #	н	OCH <sub>3</sub>	*	H	н	н	-осн <sub>2</sub> сн <sub>2</sub> -			.•
166 SO H	Ħ	осн <sub>3</sub>	н	H	N	н	-осн <sup>2</sup> сн <sup>2</sup> сн <sup>3</sup> -	•		

#### Identifying data for compounds of the invention

# NMR-data of the compounds in Table 2 (90 MHz)

Example No.	NMR-data: &(COCl <sub>3</sub> ) ppm
17	2.3(s,3H), 2.35(d,6H), 2.5(s,3H), 2.55(s,3H), 4.4(s,2H), 4.25-4.4(d,2H), 5.2-5.6(m,2H), 5.9-6.4(m,1H), 6.9(s,1H), 8.35(s,1H).
25	
27	2.2(s,3H), 2.3(s,3H), 2.6(s,3H), 4.35-4.45(d,2H), 4.45(s,2H), 5.2-5.6(m,2H), 5.85-6.35(m,1H), 6.9-7.55(m,3H), 8.3(s,1H).
29	2.2(s,3H), 2.25(s,3H), 2.4(s,3H), 4.2-4.35(c,2H), 4.4(s,2H), 5.5-5.6(m,2H), 5.85-6.3(m,1H), 6.9-7.1(d,1H), 7.3-7.55(t,2H), 8.3(s,1H).
36	1.8(m,4H), 2.75(m,4H), 3.8(s,3H), 4.25(s,2H), 5.85(m,1H), 7.05(s,2H), 7.4(d,1H), 8.3(s,1H).
<b>37</b>	1.7(m,4H), 2.3-2.7(m,4H), 3.85(s,3H), 4.6(d.2H), 6.8(s.1H), 7.05(s,2H), 7.6(m,1H), 8.3(s,1H).
44	1.2-2.0(m,10H), 2.25(s,3H), 2.3(s,3H), 2.6(m,1H), 3.75(s,3H), 4.45(s,2H), 7.1(q,1H), 7.5(m,2H), 8.35(s,1H).
56	

Evample	
No.	NMR-data: S(CDCl <sub>3</sub> ) ppm
3	2.3(s,6H), 2.35(s,3H), 2.5(s,3H), 3.75(s,3H),
	4.4(s,ZH), 7.05-7.2(d,1H), 7.25(s,1H).
	8.2-8.45(d.1H).
57	2.2(s,3H), 2.25(s,3H), 2.3(s,3H), 2.5(s,3H),
	3.45(s,3H), 3.75(s,3H), 3.85(m,4H), 4.3(s,2H),
	7.2(br.s., 1H), 8.3(s.1H).
59	2.3(s,6H), 2.4(s,3H), 2.55(s,3H), 3.5(s,3H),
	3.9(m,4H), 4.3(s,2H), 7.2(s,1H), 7.3(s,1H),
	0.4(s,lH), 9.3(br.s., lH).
66	1.2(t,3H), 2.15(s,3H), 2.2(s,3H), 2.3(s,3H),
	'2.4(s,3H), 2.8(q,2H), 3.65(s,3H), 4.8(s,2H),
	7.3(s,1H), 8.25(s,1H).
69	1.1(t,3H), 2.2(s,3H), 2.4(s,3H), 2.55(s,3H),
	2.75(q,2H), 3.85(s,3H), 4.35(s,2H), 6.75(d,1H),
	7.25(s,1H), 8.4(d,1H).
78	1.2(d,3H), 1.6(m,6H), 2.25(s,3H), 2.3(s,3H),
	3.0(m,1H), 3.75(s,3H), 4.15(m,1H), 4.45(s,2H),
	4.55(m,1H), 7.3(q,1H), 7.6(m,2H), 8.3(s,1H).
79	1.25(d,3H), 1.65(m,6H), 2.15(s,3H), 2.2(s,3H),
1	3.1(m,1H), 3.65(s,3H), 4.1(m,1H), 4.6(m,1H),
	4.8(s,2H), 7.4(q,1H), 7.7(d,1H), 7.8(s,1H),
	8.3(s,1H).
82	2.2(s,3H), 2.3(s,3H), 3.7(s,3H), 4.75(s,2H).
	7.3-8.5(m,8H).

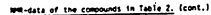
#### ------- of the compounds in Table 2. icont.

5.0-014	
··= ·	NAR-data: d(COC13) ppm
25	1.55(m,4H), 2.2(s,3H), 2.25(s,3H), 2.7-3.1(m,4H),
	3.75(s.3H), 4.35(s,2H), 6.9(d;1H), 7.3(d,1H),
	5.25(s.1H).
6	2.2(s,3H), 2.35(s,3H), 3.8(s,3H), 4.15(s,3H),
,	4.75(s,2H), 6.1(s,2H), 7.3(s,1H), 7.5(s,1H).
,	d-15(s,1H).
7	2.15(s,3H), 2.2(s,3H), 3.7(s,3H), 4.7(s,2H),
	6.05(s,2H), 7.0-7.6(m,7H), 8.15(s.1H), 8.3(s,1H).
90	2.25(s.3H), 2.1-2.4(m.2H), 2.3(s,3H), 3.75(s,3H),
30	4.2(t.4H), 4.4(s,2H), 8.75-7.2(m,5H), 7.2-7.5(n,3H),
	8.35(s.1H).
,	
92	0.7-2.05(m,13H), 2.25(s,3H), 2.3(s,3H), 2.35(s,3H),
ŀ	2.5(s,3H), 3.65-3.9(m,2H), 3.75(s,3H), 4.35(s,2H),
	7.2(s,lH), 8.3(s,lH).
93	1.25(t,3H), 2.25(s,3H), 2.3(s,3H), 2.8(q,2H),
	4.4(d,2H), 4.45(s,2H), 5.2-5.65(m,2H), 5.85-6.3(m,1H)
	7.0-7.65(m,2H), 7.5(s,1H), 8.35(s,1H).
95	0.9(s,3H), 1.0(s,3H), 1.5-1.95(m,2H), 2.15-2.45(m,]H]
i	2.25(s,3H), 2.3(s,3H), 3.7-4.0(t,2H), 3.85(s,3H),
	4.45(s.2H), 2.8-7.0(m,1H), 7.15(d.1H), 7.45-7.55
	(d,lH), 8-3(s,lH).
4.5	2.25(s,3H), 2.40(s,3H), 3.8 and 3.85(2s, total 3H),
	3.80(s,3H), 4.8 and 4.85(2s,totel 2H), 8.35-7.95
	(m,8H), 8.35(s,1H).

o

The state of the s

Example	
No.	NMR-data: #(CDC1 <sub>3</sub> ) ppm
103	2.3(s,3H), 2.35(s,3H), 3.0(t,2H), 3.35(s,3H),
	3.65(t,2H), 3.8(s,3H), 4.4(s,2H), 6.8-7.6(m,4H),
	8.25(s,1H).
107-108	2.2(s,3H), 2.35(s,3H), 3.75(s,3H), 3.9 and 3.95
	(2s,total 3H), 4.15(s,3H), 4.75(s,2H), 7.07-7.95
	(=,3H), 8.15(s,1H).
102	1.32(s,9H), 2.08(s,3H), 2.15(s,3H), 4.09(d,2H),
	4.74(s,2H), 5.10-5.45(m,2H), 5.73-6.25(m,1H),
	7.28-7.73(m,3H), 8.27(s,1H).
139	2.22(s,3H), 2.29(s,3H), 3.75(s,3H), 4.40(s,2H),
	7.38-7.58(m, 1H), 7.87-8.02(m, 2H), 8.29-8.47(m, 1H),
	8.70-9.00(m,2H).
110	1.25(d,6H), 1.6-2.15(m,4H), 2.25(s,3H), 2.3(s,3H),
	3.0(m,1H), 3.7-4.05(m,4H), 4.25(m,1H), 4.5(s,2H),
	7.15(q,1H), 7.5(s,1H), 7.55(d,1H), 8.3(s,1H).
111	1.3(d,6H), 1.55-2.15(m,4H), 2.2(s,3H), 2.25(s,3H),
	3.05(m,1H), 3.65(d,2H), 3.9(m,2H), 4.2(m,1H), 4.8
	(s,2H), 7.3(d,1H), 7.4-7.8(m,2H), 8.3(s,1H).
119	2.3(s,3H), 2.35(s,3H), 3.15(t,2H), 3.7(s,3H),
	4.25(t,2H), 4.4(s,2H), 6.9(q,1H), 7.15(d,1H), 7.3-
	7.6(m,6H), 8.35(s,1H).
 125	2.3(s,3H), 2.35(s,3H), 2.8(s,3H), 3.8(s,3H), 4.5
	(s,2H), 7.5(d,1H), 7.75(d,1H), 8.05(s,1H), 8.4(s,1H).



Example	NMR-data: d(CDC1 <sub>3</sub> ) ppm
No.	The state of the s
126	2.2(s,6H), 2.8(s,3H), 3.7(s,3H), 4.85(s,2H), 7.6
	(q.1H), 7.85(d.1H), 8.15(s.1H), 8.25(s.1H).
127	2.25(d,6H), 3.75(s,3H), 4.9(d,2H), 7.8(d,1H),
	8.3(s,1H), 8.3(q,1H), 8.65(d,1H).
134	2.2(d,6H), 2.35(d,6H), 3:1(s,6H), 3.7(s,3H), 4.95
	(s,2H), 7.2(s,1H), 7.6(s,1H), 8.3(s,1H).
112	2.1(s,3H), 2.25(s,3H), 2.3(s,3H), 2.65-3.2(m,4H).
	4.4(d.2H), 4.42(s,2H), 5.2-5.6(m,2H), 5.9-6.4(m,1H),
	7.1(dd,1H), 7.4(d,1H), 7.5(d,1H), 8.35(s,1H).
121	2.25(s,3H), 2.35(s,3H), 3.8(s,3H), 4.45(s,2H),
	7.45-8.0(m,7H), 8.15(s.1H), 8.4(s.1H).
122	2.2(s,6H), 3.7(s,3H), 4.8(d,2H), 7.5-8.05(m,7H),
	8.2(s,1H), 8.25(s,1H).
144	2.25(s,3H), 2.35(s,6H), 2.38(s,3H), 2.55(s,3H),
	4.4(s,2H), 7.15(d,1H), 7.3(s,1H), 8.4(d,1H).
145	2.15(s.3H), 2.23(s.3H), 2.27(s.3H), 2.4(s.3H),
	2.47(s.3H), 4.8(s.2H), 7.1(d.1H), 7.3(s.1H),
	8.37(d,1H).
151	2.2(s,3H), 2.23(s,3H), 2.35(s,3H), 2.4(s,3H),
· '	2.47(s.3H), 4.8(d.2H), 7.0(s.1H), 7.1(d.1H), 8.37
	(d,1H).
130	3.85(s,3H), 4.65(s,2H), 6.8-7.8(#,7H), 8.55(d,1H)

NMR-data of the compounds in Table 2. (cont.)

Example No.	MMR-data: &(CDCl <sub>3</sub> ) ppm
131	3.85(s.3H), 4.95(d.2H), 6.65-7.60(m.7H), 8.45(d.1H).
160	1.15(t,3H), 2.20(s,3H), 2.27(s,3H), 2.49-2.73(m,2H), 2.89-3.13(m,2H), 3.72(s,3H), 4.09(q,2H), 4.37(s,2H), 6.98 and 7.08(dd,1H), 7.30-7.55(m,2H), 8.28(s,1H).
154	1.1-2.1(m,13H),2.3(s,3H),2.5-2.8(m,3H), 4.4(s,2H), 7.1-7.65(m,4H), 8.5(s,1H)
156	1.1-2.0(m,11H), 2.25(s,3H), 2.3(s,3H), 3.45(s,3H), 3.7(t,2H), 4.0(t,2H), 4.4(s,2H), 7.05-7.65(m,3H), 8.35(s,1H)
164 (270 MHz)	2.13(m,2H),2.88(t,2H),3.82(s,3H),4.26(t,2H), 4.69(s,2H),6.7-6.85(m,2H),7.04(d,1H), 7.39(d,1H),8.1(d,1H).



Preparation of intermediates

Example 11. Method A. Preparation of 4,5,7trimethyl-2-mercapto-1H-benzimidazole.

2-Nitro-3,4,6-trimethylaniline (10.2 g, 0.057 mol) was dissolved in 95% ethanol (900 ml) and hydrogenated in the presence of Pd/C-catalyst until the theoretical amount of hydrogen had been consumed (1 hour). The whole mixture was transferred to another flask and potassium ethylxanthate (12.8 g,

10 0.080 mol) dissolved in water (12.5 ml) was added. The mixture was refluxed overnight, 2M NaOH (20 ml) was added and the volatiles were evaporated off. The residue was dissolved in methanol (300 ml) and the catalyst was filtered off. Part of the solvent (200

15 ml) was evaporated off. Water (100 ml) was added and the mixture was acidified with acetic acid (10 ml) dissolved in water (20 ml). The crystalline precipitate was filtered off, washed with water and dried under reduced pressure, giving the desired product (7.2 g.

20 66%), NMR: δ(COCl<sub>3</sub>) 2.0(s,3H), 2.05(s,3H), 2.1(s,3H), 3.3(br.s,1H), 6.5(s,1H).

Example 12. Method B. Preparation of 4,6,7 - trimethyl - 5 - methoxy - 2 - mercapto - 1H - benzimidazole.

25 A solution of 4 - methoxy - 3,5,6 - trimethyl - 1,2 - phenylenediamine (1.8 g, 0.010 mol) and triethylamine (2.1 g), 0.021 mol) in CHCl<sub>3</sub> (15 ml) was added dropwise to a stirred solution of thiophosgene (0.60 g, 0.0052 mol) in CHCl<sub>3</sub> (5 ml). The mixture was then 30 stirred at room temperature for 1 hour. Water (15 ml) and triethylamine (0.5 g) was added and the mixture was stirred for 1 hour. The precipitate was filtered off,

washed with water and dried in the air giving the

desired product (0.96 g, 43%), NMR:  $\delta(COCl_3)$ 

35 2.5(s,3H), 2.65(s,6H), 3.65(s,3H), 12.0(br.s.,1H). Example 13. Method C. Preparation of 4-allyloxy - 3,5 dimethyl - 2 - pyridinyl - methanol.

4 - Allyloxy - 2,3,5 - trimethyl - pyridine N-oxide (4.0 g, 0.021 mol) was added dropwise under stirring to acetic anhydride (8.0 ml, 0.062 mol) preheated to 80°C, giving a final temperature of 120°C. The mixture was then heated at 80°C for 1 hour. Methanol (15.0 ml) was added and the mixture was kept at 80°C for 15 min. The volatiles were evaporated under reduced

pressure. 10% HCI (20ml) was added and the mixture was heated at 90°C for 1 hour and then cooled to room temperature. Excess 2M NaOH was added and the mixture was extracted with CH₂Cl₂. The organic phase was separated out and dried. Volatiles were evaporated off giving the desired product as an oil

evaporated off giving the desired product as an oil (3.0 g, 75%), NMR: δ(COCI<sub>3</sub>) 2.1(s,3H), 2.25(s,3H), 4.4(m,2H), 4.65(s,2H), 4.75(s, 1H), 5.2-5.65(m,2H), 5.9-6.45(m,1H), 8.3(s,1H).

Example 14. Method D. Preparation of 4 - allyloxy - 3,5
55 - dimethyl - 2 - pyridinyl - methyl chloride hydrochloride.

Thionyl chloride (4.0 ml) dissolved in CH<sub>2</sub>Cl<sub>2</sub> (12 ml) was added dropwise to a stirred solution of 4-allyloxy-3,5-dimethyl-2-pyridinylmethanol (8.0 g, 0.041 mol) in CH<sub>2</sub>Cl<sub>2</sub> (50 ml), maintaining the temperature below 6°C. Then the mixture was stirred at room temperature for 45 min (final temperature 15°C). Isopropanol (2 ml) was added and the solution was heated shortly at 35°C. The solvent was evaporated off and the crystalline residue was recrystallized from ethanol/ether giving the desired product (3.0 g, 29%), m.p. 115°C.

Table 3a. Intermediates. Summary of working examples.

$$\begin{array}{c|c}
R^{2a} & & \\
R^{3a} & & & \\
R^{3a} & & & \\
R^{4a} & & & \\
R^{5a}
\end{array}$$

No.	Z <sup>la</sup>	Rla	R <sup>2a</sup>	R <sup>3a</sup>	R <sup>4a</sup>	R <sup>Sa</sup>	Method <sup>x).</sup> (Ex. No.		Mp (OC) other data
15	SH	о,	CH <sub>3</sub>	CH3	сч3	н	A(Ex 11)	19	HPR
16	KZ	CH <sub>3</sub>	сн <sub>3</sub>	CH3	н	H	A(Ex 11)	66	10%
11	SH	CH3	CH <sub>3</sub>	H	СНЗ	н	A(Ex 11)	66	NetR
17	SH	н	$\prec \frown$	н	н	н	A(Ex 11)	71	NIR
18	SH	CHa	OCH <sub>3</sub>	СНЗ	н	н	A(Ex 11)	78	NMR
19	SH	CH3	OCH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>	CH <sub>3</sub>	н	н	A(Ex [3)	. 85	NPER
110	SH	CH <sup>3</sup>	C <sub>2</sub> H <sub>S</sub>	CH3	н	H	A(Ex 11)	89	NHR
111	SH	٠.	0CH2CH2CH20-(O)	н	н	н	A(Ex 11)	14	167
112	SH	٠.	O(CH <sub>2</sub> )6CH <sub>3</sub>	CH3	н	H	A(Ex 11)	73	KX
12	SH	СН,	ОСН	CH3	Сн3	н	B(Ex 12)	43	NMR
113	SH	,	•сн-сн-сн-сн <sub>2</sub> сн <sub>2</sub> -		н	н	A(Ex 11)	23	たる

a) Method A: The 1,2-phenylenediamine is reacted with C2M50C52K Method B: The 1,2-phenylenediamine is reacted with CSC12

very of working examples.

No.	2 <sup>2</sup> a	R <sup>64</sup>	g <sup>7</sup> a	R <sup>Sa</sup>	Salt/Base	Method XX) (Ex. No.)	Yield (%)	Mp (°C) other data
13	ОН	CH.	OCH <sub>2</sub> CH+CH <sub>2</sub>	CHa	Base -	C(Ex 13)	75	NAC
14	CI	-	0CH <sub>2</sub> CH+CH <sub>2</sub>	CH3	HC1	D(Ex 14)	29	1150
[14		CH3	OCH <sub>2</sub> C=CH	CH <sub>2</sub>	Base	C(Ex 13)	88	70 <sup>0</sup>
115		CH <sub>2</sub>	OCH <sub>2</sub> C±CH	CH <sub>3</sub>	нс1	0(Ex 14)	76	1350
116	ОН	H	-(CH <sub>2</sub> )	•	Base	C(Ex 13)	35	184R
117	C1	н	-(CH <sub>2</sub> )		нс1	D(Ex 14)	72	NHR .
	OH	сн,	OCH_CH_CH(CH		Base	C(Ex 13)	51	HIR
119		•	OCH_CH_CH(CH		нс1	D(Ex 14)	95	
120	OH		осн <del>,</del>	CH,		C(Ex 13)	30	MAR
:21	Ç1	•	осн; <del>(</del> )	CH.		D(Ex 14)	62	133
:22	ύн	CH3	ענ <sub>י</sub> אל	CH <sub>3</sub>	Base	C(Ex 13)	70	8.p. 120- 26 C/0.4
123	C1	CH,	0C <sub>2</sub> H <sub>5</sub>	CH.	HC1	D(Ex 14)	89	157
124		•	-CH-0-	н .	Base	C(Ex 13)	18	JH MAK.
125	c1	-CH:	-CH-0-	н	HC1	D(Ex 14)	95	195

 $^{
m XX)}$ Method C: Rearrangement of the pyridine N-oxide with  ${
m (CH_3CO)}_2{
m O}$ . Method D: Chlorination with SOC12.

NMR-	-data of the compounds in Table 3a and	Table
3b	•	

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NMR-data: ठ(ppm)

δ(DMSO-d<sub>6</sub>) 2.05(s,6H), 2.2(s,6H). 5 15

δ(CDCl<sub>3</sub>) 2.05(s,3H), 2.15(s,3H), 2.2(s,3H, 16 3.2(s,2H), 6.7(s,1H).

δ(CDCl<sub>3</sub>) 2.0(s,3H), 2.05(s,3H), 2.1(s,3H), 3.3(br.s.,1H), 6.5(s,1H).

δ(DMSO-d<sub>6</sub>) 1.1-2.05(m,10H), 2.4(m,1H), 10 17 6.85-7.05(m,3H).

δ(DMSO-d<sub>6</sub>) 1.95(s,3H), 2.0(s,3H), 3.35(s,3H), 6.55(s,1H).

δ(CDCl<sub>3</sub>) 2.1(s,3H), 2.15(s,3H), 3.2(s,3H),

3.35-3.8(m,4H), 6.6(s,1H). 15

110 δ(CDCl<sub>3</sub>+DMSO-d<sub>6</sub>) 1.05(t,3H), 2.3(s,3H), 2.35(s,3H),

2.6(q,2H), 6.85(s,1H).

112 δ(CDCl<sub>2</sub>) 0.5-1.7(m,13H), 2.0(s,3H), 2.1(s,3H),

3.15(s,2H), 3.35-3.6(m,2H), 6.6(s,1H).

δ(CDCl<sub>3</sub>) 2.5(s,3H), 2.65(s,6H), 3.65(s,3H), 12.0(br.s.,1H).

113 δ(CDCI<sub>2</sub>) 3.35(s,2H), 3.4(s,2H), 7.15-8.05(m,4H),

12.65(br.s.,1H), 13.3(br.s.,1H). δ(CDCl<sub>3</sub>) 2.1(s,3H), 2.25(s,3H), 4.4(m,2H),

4.65(s,2H), 4.75(s,1H), 5.2-5.65(m,2H), 5.9-6.45(m,1H), 8.3(s,1H).

116 δ(CDCl<sub>3</sub>) 1.5-1.9(m,4H), 2.5-2.8(m,4H), 4.7(s,2H), 7.3(s,1H), 8.2(s,1H).

30 117 118 δ(CDCl<sub>3</sub>) 1.0(s,3H), 1.05(s,3H), 1.5-2.05(m,3H), 2.15(s,3H), 2.3(s,3H), 3.75-4.0(t,2H),

4.15-4.5(br.s.,1H), 4.65(s,2H), 8.3(s,1H).

120 δ(CDCl<sub>3</sub>) 1.7-2.2(m,4H), 2.15(s,3H), 2.25(s,3H), 3.75- 4.05(m,4H), 4.15-4.4(m,1H), 4.6(s,2H),

35 8.25(s,1H).

124 δ(CDCl<sub>3</sub>) 8.55(d,1H), 7.8(d,1H), 7.5(d,1H), 7.0(d,1H), 5.1(s,2H).

Pharmaceutical preparations containing a compound 40 of the invention as active ingredient are illustrated in the following examples.

Example 167. Syrup

A syrup containing 1% (weight per volume) of active substance was prepared from the following

45 ingredients:

4,6-Dimethyl-5-ethyl-2-[[(4-methoxy-3.5-dimethyl-2-pyridinyl)methyl]thio]-

1.0 g 1H-benzimidazole-HCI 30.0 g Sugar, powder 0.6 g 50 Saccharine 5.0 g Glyceroi 0.059 Flavouring agent 5.0 q Ethanol 96% Distilled water q.s. to a final volume of 100 ml

Sugar and saccharine were dissolved in 60 g of warm water. After cooling the acid addition salt was dissolved in the sugar solution and glycerol and a solution of flavouring agents dissolved in ethanol

60 were added. The mixture was diluted with water to a final volume of 100 ml.

The above given active substance may be replaced with other pharmaceutically acceptable acid addition salts.

Example 168. Enteric-coated tablets

An enteric-coated tablet containing 20 mg of active compound was prepared from the following ingredients: 1 5,6-Methylenedioxy-2-[[(4-methoxy-

5		3,5-dimethyl-2-pyridinyl)methyl[sulfinyl]-	
•		1H-benzimidazole	200 g
		Lactose	700 g
		Methyl cellulose	6 g
		Polyvinylpyrrolidone cross-linked	50 g
10		Magnesium stearate	15 g
10		Sodium carbonate	6 g
		Distilled water	q.s.
	11	Cellulose acetate phthalate	200 g
	••	Cetyl alcohol	15 g
15		Isopropanol	2000 g
13		Methylene chloride	2000 g
	1	5,6 - Methylenedioxy - 2 - [[(4 - methoxy	- 3,5 -

dimethyl - 2 - pyridinyl)methyl]sulfinyl] - 1H benzimidazole, powder, was mixed with lactose and 20 granulated with a water solution of methyl cellulose and sodium carbonate. The wet mass was forced through a sieve and the granulate dried in an oven. After drying the granulate was mixed with polyvinylpyrrolidone and magnesium stearate. The dry mix-

25 ture was pressed into tabled cores (10 000 tablets), each tablet containing 20 mg of active substance, in a tabletting machine using 6 mm diameter punches. II A solution of cellulose acetate phthalate and cetyl

alcohol in isopropanol/methylene chloride was 30 sprayed onto the tablets I in an Accela Cota, Manesty (RTM) coating equipment. A final tablet weight of 110 mg was obtained.

Example 169. Solution for intravenous administra-35 tion

A parenteral formulation for intravenous use, containing 4 mg of active compound per ml, was prepared from the following ingredients: 4,6-Dimethyl-5-ethyl-2-[[(4-methoxy-

40 3,5-dimethyl-2-pyridinyl)methyl]thio]-4 g 1H-benzimidazole Polyethylene glycol 400 for injection 400 g q.s. Disodium hydrogen phosphate 1000 ml Sterile water to a final volume of

4,6-Dimethyl-5-ethyl-2-[[(4-methoxy-3,5dimethyl-2-pyridinyl)methyl]thio]-1H-benzimidazole was dissolved in polyethylene glycol 400 and 550 ml of water was added. pH of the solution was

50 brought to pH 7.4 by adding a water solution of disodium hydrogen phosphate and water was added to a final volume of 1000 ml. The solution was filtered through a 0.22 µm filter and immediately dispensed into 10 ml sterile ampoules. The ampoules were

55 sealed.

45

Biological tests I. Inhibiting effect in vitro on acid secretion in isolated rabbit gastric glands Test Method

60 Gastric gland preparation

Isolated rabbit gastric glands were prepared as described by Berglindh et al., Acta physiol. scand. 1976. 96. 150-159. This method involves vascular perfusion of the raibbit stomach via the gastric 65 arteries, scraping and scissor mincing of the sepa-

rated gastric mucosa and collagenase (0.1%, Type I, Sigma Chemicals, St. Louis, MO. USA) digestion at 37°C for 60-90 min. The glands are then harvested and filtered through nylon cloth to remove coarse frag-70 ments. The glands are thereafter incubated at 37°C in a medium containing NaCl 132.4 mM, KCl 5.4 mM, NaH<sub>2</sub>PO<sub>4</sub>, 5.0 mM, NaH<sub>2</sub>PO<sub>4</sub>, 1.0 mM, MgSO<sub>4</sub> 1.2 mM, CaCl<sub>2</sub> 1.0 mM, glucose 10 mM, and 1 mg/ml rabbit albumine, pH 7.4.

75 Measurement of acid secretion

The acid secretion in the isolated gland preparation was recorded by measuring the uptake of 14C-labelled aminopyrine into the glands as described by Berglindh et al., Acta physiol. scand. 1976. 97. 401-414. 80 Accumulation of aminopyrine in the glands indicates gastric acid secretion within the glands. The standard medium contained 10-6M 14C-aminopyrine (Amersham, Great Britain). After the incubation period, the glands were centrifuged, the supernatant was re-85 moved and the glands dried, weighed and dissolved in Soluene -350 (Packard, IU. USA). Samples of the supernatant and glands were separately counted in a scintillation counter. The accumulation of 14C-labelled aminopyrine in the glands was calculated as detailed by Berglindh et al., Acta physiol. scand. 1976.

Experimental protocol

Glands were incubated for 60 min. in the presence of  $5 \times 10^{-5}$ M histamine and the test compound to be studied. The free base of the test compound was dissolved in methanol. The final concentration of methanol was 1% in the incubation medium, having no influence on the aminopyrine accumulation ratio. For each test compound a complete dose-response

100 curve was generated by testing doses in duplicate in the concentration range 10<sup>-7</sup>M to 10<sup>-4</sup>M. The logarithm of the concentration (in M) of the test compounds giving 50% inhibition of the aminopyrine accumulation in the glands (IC50) is listed in Table 4 105 below.

II. Inhibiting effect in vivo on gastric acid secretion in conscious dog

Test Method

Chronic gastric fistula dogs were used. These dogs 110 have been surgically provided with a gastric cannula in the stomach and a duodenal fistula used for direct introduodenal administration of test compounds. Following a 4 weeks' recovery period after surgery. tests were performed once a week on each dog. Food 115 and water were withdrawn 18 hours before each test.

Gastric acid secretion was induced by continuous infusion of histamine at individual doses 1100-300 nmol/kg, h), resulting in submaximal secretion of gastric acid. At least 2 hours after onset of stimula-

120 tion, when the gastric acid secretion had reached a steady level, the test compounds in the form of free base suspended in 0.5% Methocel (RTM) (90 HG. 15.000, Dow Chem. Corp.), were given intraduodenally at doses from 1 to 8 µmoVkg. The gastric juice was

125 collected by free flow from the gastric cannula in consecutive 30 minutes samples for 3 hours. The samples were titrated to pH 7.0 with 0.1 M NaOH using a Radiometer automatic titrator and the acid output was calculated.

The per cent inhibition of acid secretion was

calculated by comparing in each dog the acid output in the tests to the acid output in control tests when

only the vehicle was given. The peak inhibitory effect for each compound is given in Table 5 below.

Table 4 Biological effects in isolated rabbit gastric glands

<b>Ja.</b>	1	R13	z1	R <sup>2</sup>	R <sup>3</sup>	R.	R <sup>5</sup>	26	R <sup>7</sup>	2.5	-low IC50
12	- 50	11	CH,	CH	α,	ca,		CH <sub>3</sub>	осн <sub>3</sub>	CH.)	6-5
. 10	50	•	CK,	CH <sub>2</sub>	CH <sub>3</sub>	1	×	CH3	осн3	CH.	6.5
· 37	\$0	1	1	оси		*	ĸ	×	-(C	H <sub>2</sub> )4=	5.0
4.5	50		1	ocs,cs		11		CH <sub>3</sub>	oca,	CX3	4.4.
əl	50	ı		CH_OH ·	CIL <sub>3</sub>		1	CH <sub>3</sub>	оси3	CH.	6.1
104	50	1		CH <sub>2</sub> CH <sub>2</sub> OCH <sub>3</sub>		1	Ħ	CN3	OCH <sub>3</sub>	CIL)	5-7
	50		CIL,	ocu,	CII,	*	×	CH3	OCH <sub>3</sub>	CH <sub>3</sub>	6.5
1	30	=	ац	OCH	CH3	19	Ħ	CH <sup>3</sup>	CN,	Ħ	6,7
34	\$0	ъ	CIL,	003,01,001,	CH <sub>3</sub>		ĸ	CH3	OCH <sub>3</sub>	CH3	5.9
₩	\$0		ai,	001,01,001,	α,		1	Ħ	CM	CH3	5.4
62	so	<b>.</b>	a,	cocar,	CH <sub>3</sub>		1	CH <sub>3</sub>	ося,	CH <sub>3</sub>	6-2
64	so		СВ,	cocar,	CN,	*		CT <sub>3</sub>	19	CH <sub>3</sub>	5.8
••	50		α,	coc,s,	CH,	H		CH,	осн,	CH,	6.0
	-		,	• •	•				•		Cont.

•	06	E	

coat								- 6		. 8	
No.	x	R13	* * ·	R <sup>4</sup>	R"	R.	R <sup>3</sup>	R*	1'	R <sup>8</sup>	-log IC50
64	<b>50</b>	×	95	C2H2	CH <sub>3</sub>	Ħ,	я	СНЗ	скэ	CH3	6.5
ν	50	×	(P)	c <sub>2</sub> n <sub>5</sub>	CH3	H	Ħ	CH3	осн3	И	5.9
72	so	×	c <sub>2</sub> A	· Car	c2×2	H	Ħ	CN,	осн3	CH3	5.0
74	so	1	сн <sub>3</sub> \	оси <sub>з</sub> .	CH3	ᅄ	Ħ	CH3	осн <sub>3</sub>	CH3	6.2
79	so	<b>T</b>	*	25	я	H	Ħ	CH <sub>3</sub>	осн 3	CH3	5.0 .
81	so	ı	2	<b>/</b> оси <sub></sub> 0-		Ħ	Ħ	СНЭ	осн3	CH3	6.1
<b>#3</b>	50	Ħ	-CH	-ся-ся-фя-	H	H	Ħ	СĸЭ	∞сн <sub>3</sub>	CH,	{5.5 5.3
107	so	H	n	OCH <sub>3</sub>	×	H	COZCH	Сн3	осн3	CH3	<b>5.8</b>
108	so		M	in .	осн <sub>3</sub>	H	CO <sub>2</sub> CH <sub>3</sub>	CH3	осн,	CH3	

cont.

coal.

Ho.	- x	R15	A <sup>1</sup>	R <sup>2</sup>	<b>8</b> 3	R <sup>4</sup>	R <sup>5</sup>	. g*	R)	R <sup>6</sup> log	1C 20
10	so	и	CH,	CH,	си <sub>3</sub>	CH,	н	CH3	OCH2CH-CH2	CN <sub>2</sub>	6.1
14	50	M	•	cx,	CH <sub>3</sub>	N			OCH2CH-CH2		6.1
18	so	8	•	CH <sub>3</sub>	ĸ	СНЗ	M	CH <sub>3</sub>	OCH2CH-CH2	CH <sub>3</sub>	5.9 .
20	50		•	cn,	Ħ	CH <sub>3</sub>	Ħ	CH.	och <sub>3</sub>	CH3	6.0
22	so	· M	•	CH3	н	н	H	CH,	oca <sub>2</sub> ca-ca <sub>2</sub>	ca <sub>3</sub>	6.0
24	so	1	-	CH,	cH <sub>3</sub>	Ħ	×	CH <sub>3</sub>	OCH 2CH-CH2	CH <sub>3</sub>	6.0
26	. 50	a	CH,	н	н	CH <sub>3</sub>	Ħ	CII3	OCH2CH-CH2	cx,	5.9
28	50	W	CH3		H	н	Ħ	CH <sub>3</sub>	OCH2CH-CH2	CH <sup>3</sup>	5.9
30	50	al	•	си,	н	н	Ħ	CH3	0CH2CH-CH2	cx3	5.9.
32		Ħ		OCN,	н	Ħ	¥	CH <sub>3</sub>	0CH2CH-CH2	CH,	5.6
34	so	н		OCH,	ĸ	`N	H	CH3	OCH <sub>2</sub> CFCH	CH <sub>3</sub>	5.0
35	so	11	M	осн	. 11	H	Ħ	Ħ	ocat <sub>3</sub>	. c <sub>2</sub> # <sub>5</sub>	5.6
41		н	CH3	•	СНЗ	d	H	CH3	OCH <sub>2</sub> CH-CH <sub>2</sub>	CH <sub>3</sub>	5.9
45			,	_	н	Ħ		_	ocii,		6.1
_	_	٠.	•	_	,			_			•

cont.

No.	<u>x</u>	RIS	Rì	R <sup>2</sup>	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R	R <sup>2</sup>	R <sup>8</sup>	-10E IC50
 5>	50	`		COCE,	Ch <sub>3</sub>	н	М	CH3	OCH <sub>2</sub> CH-CH <sub>2</sub>	CII.)	5.3
67	so	h	-CH.	CH2CH2CH2-	H	Ħ	h	CH3	och <sub>3</sub>	CH.3	6.3
91	>0		h .	och,ch,ch,o-©	h		H	CH <sub>3</sub>	OCH3	CH <sup>3</sup>	5.8
. 2	so	H	Cit <sub>2</sub>	0(Ch <sub>2</sub> ),Ch <sub>3</sub>	CH3	H	×	CH.	оси 3	CH.	5.9
94	so			C,n,	h	h	h	CH3	оси <sub>2</sub> си-си <sub>2</sub>	CH <sup>3</sup>	4.6
76	SU	<b>h</b>		oca <sub>3</sub>	h	h	H	CH <sup>3</sup>	001201201013	2 <sup>CH</sup> 3	6.1
78	so		-0	I-CH-CH-CGH <sub>2</sub> CH <sub>2</sub> -		н	H	CH3	осн3	CH3	5.6
102	sc	н	н	C(CH <sub>3</sub> ) <sub>3</sub>	н	н	н	CH3	OCH <sup>2</sup> CH-CH <sup>2</sup>	CH3	5.9
104	so	н	×	CH2CH2OCH3	H	н :	н	CH3	OCH3	CH <sup>3</sup>	5.7
106	so	H	¥	٠,	<b>`</b> 0-	н	н	CH3	осн <sub>3</sub>	CH3	6.0
111	so	н	н	сн(сн <sub>3</sub> ) <sub>2</sub>	н	H	H	CH3	OCH TO	CH3	6.2
113	50		н	CH <sub>2</sub> CH <sub>2</sub> COCH <sub>3</sub>	н	н	н	CH <sup>3</sup>	OCH <sup>2</sup> CH=CH <sup>2</sup>	CH3	5.8
118	so		н	<b>~</b> ⊚	H	H	×	CH3	_	CH <sub>3</sub>	6.4

ont.

cont.

mo.	1	R <sup>15</sup>	R <sup>1</sup>	R <sup>2</sup>	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup> .	R <sup>8</sup>	-log IC <sub>50</sub>
120	so	н	н	OCH <sub>2</sub> CH <sub>2</sub> -O)	н	н	н	СНЭ	осн3	CH3	6.3
124	SO.	H	н	-⊚ ·	H	H	М	CH3	осн <sub>3</sub>	СНЗ	7.0
129	so	н	н	Br .	н .	н	н.	CH3	OCH <sup>2</sup> CH-CH <sup>2</sup>	CH3	
142	so	×	×	-OCH	20-	H	H	CH3	CH <sub>3</sub>	Сн3	6.0
143	50	н	H	POCH,	CH3	н	н	H	осн <sub>3</sub>	CZHS	6.1
145	so	н	CH,	он,	CH <sub>3</sub>	H	H	CH <sub>3</sub>	CH <sub>3</sub>	н	6.2
147	so	H	•	CH <sub>3</sub>	CH <sub>3</sub>	н	ж	H	CH3	CH3	6.4
149	so	н	•	CH <sub>3</sub>	CH3	H	H	CH3	H	CH3	6.2
151	so	н	CH	CH	H	CH3	H	CH <sub>3</sub>	CH3	н	6.3
153	so	н	CH,	CH	CH <sub>3</sub>	н -	н	CH <sub>3</sub>	oc <sub>z</sub> H <sub>S</sub>	CH <sub>3</sub>	5.2
77	so	н	H		CH3	H	H	H	осн3	C2H5	6.0
159	š	H	H	cr <sub>3</sub>	H	Ħ	H	CE3	OCH 2	CH <sup>3</sup>	6.3

## Table 5 Biological effects in conscious dogs

								•				
No.	x	R <sup>15</sup>	R <sup>1</sup>	RZ	R <sup>3</sup>	R <sup>4</sup>	R <sup>5</sup>	R <sup>6</sup>	R <sup>7</sup>	R <sup>8</sup>	(1.D.) DOSE (pmol/kg	) % INHIB
84	s	н	н		-CH-CH-CH-CH-	н	н	CH3	осн <sub>3</sub>	CH3	8	- 85
109	s	н	×	SCH.	н	н	н	CH <sub>3</sub>	осн <sub>3</sub>	CH3	8	60

# Comment to the test results

It is seen in Table 4 and Table 5 that the tested compounds potently inhibited gastric acid secretion both in vitro and in vivo.

### 5 CLAIMS

## 1. A compound of the formula

wherein

 $R^{15}$  is H, CH<sub>3</sub> or  $C_2$ H<sub>5</sub>;

- 10 R1, R2, R3 and R4, which are the same or different, are
  - (a) H
  - (b) halogen
  - (c) -CN
  - (d) —(HO
- 15 (e) —(.F<sub>3</sub>

- 0 || (f) —C—R'
- (g) --O--C--R<sup>12</sup>
- (h) -CH(OR13)2
- (i)  $-(Z)_n A D$
- g (j) aryl
  - (k) aryloxy
  - (I) alkylthio containing 1-6 carbon atoms
  - (m) -NO<sub>2</sub>
  - (n) alkylsulfinyl containing 1-6 carbon atoms or

# 25 wherein

- (o) adjacent groups R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> together with the adjacent carbon atoms in the benzimidazole ring form a 5-, 6- or 7-membered monocyclic ring or a 9-, 10- or 11-membered bicyclic ring which rings may be
- 30 saturated or unsaturated and may contain 0-3 hetero atoms selected from—N— and—O—, and which rings may be optionally substituted with 1-4 substituents selected from alkyl groups with 1-3 carbon atoms, alkylene radicals containing 4-5 carbon atoms
- 35 giving spiro compounds, or two or four of these substituents together form one or two oxo groups.

(-C-), whereby if R1, R2, R3 and R4 together with

the adjacent carbon atoms in the benzimidazole ring form two rings they may be condensed with each other, in which formulas R11 and R12, which are the same or different, are

(a) aryl,

(b) alkoxy containing 1-4 carbon atoms,

(c) alkoxyalkoxy containing 1-3 carbon atoms in each alkoxy part,

(d) arylalkoxy containing 1-2 carbon atoms in the 10 alkoxy part,

(e) aryloxy,

(f) dialkylamino containing 1-3 carbon atoms in each alkyl residue, or

(g) pyrrolidino or piperidino, optionally substituted with alkyl containing 1-3 carbon atoms; R13 is (a) alkyl containing 1-4 carbon atoms, or

(b) alkylene containing 2-3 carbon atoms;

Zis-O-or-Cnis0or1;

A is (a) alkylene containing 1-6 carbon atoms

(b) cycloalkylene containing 3-6 carbon atoms

(c) alkenylene containing 2-6 carbon atoms

(d) cycloalkenylene containing 3-6 carbon atoms,

25 or

20

(e) alkynylene containing 2-6 carbon atoms;

Dis(a) —CN 0

(c) -(Y)<sub>m</sub>-(C),-R<sup>10</sup>

30 wherein

R<sup>9</sup> is (a) alkoxy containing 1-5 carbon atoms, or

(b) dialkylamino containing 1-3 carbon atoms in each alkyl residue;

mis0or1;

35 ris0or1;

Yis (a) -0-

(b) -NH-

(c) -NR10-:

R<sup>10</sup> is (a) H

(b) alkyl containing 1-3 carbon atoms,

(c) arylalkyl containing 1-2 carbon atoms in the alkyl part, or

(d) aryl;

R<sup>5</sup> is (a) H or

0

(b) -C-R14:

R<sup>14</sup> is (a) alkyl containing 1-6 carbon atoms,

(b) arylalkyl containing 1-2 carbon atoms in the alkyl part

(c) aryl

(d) alkoxy containing 1-4 carbon atoms

(e) arylalkoxy containing 1-2 carbon atoms in the alkyl part

(f) aryloxy

(g) amino

(h) mono- or dialkylamino containing 1-4 carbon atoms in each alkyl residue

(i) arylalkylamino containing 1-2 carbon atoms in the alkyl part

(j) arylamino;

R<sup>6</sup> and R<sup>8</sup>, which are the same or different, are

(a) Hor

(b) alkyl containing 1-5 carbon atoms;

R<sup>7</sup> is (a) H

(b) alkyl containing 1-8 carbon atoms

(c) alkoxy containing 1-8 carbon atoms

(d) alkenyloxy containing 2-5 carbon atoms

(e) alkynyloxy containing 2-5 car .n atoms

(f) aikoxyalkoxy containing 1-2 carbon atoms in

70 each alkoxy group

(g) dialkylaminoalkoxy containing 1-2 carbon atoms in each of the alkyl residues on the amino nitrogen and 1-4 carbon atoms in the alkoxy group

(h) oxacycloalkyl containing one oxygen atom and

75 3-7 carbon atoms

(i) oxacycloalkoxy containing two oxygen atoms and 4-7 carbon atoms

(j) oxacycloalkylalkyl containing one oxygen atom and 4-7 carbon atoms

(k) oxacycloalkylalkoxy containing two oxygen atoms and 4-6 carbon atoms, or

(I) R<sup>6</sup> and R<sup>7</sup>, or R<sup>7</sup> and R<sup>8</sup> together with the adjacent carbon atoms in the pyridine ring from a ring wherein the part constituted by R<sup>6</sup> and R<sup>7</sup>, or R<sup>7</sup> and

85 R<sup>8</sup>, is

-CH=CH-CH=CH-

-O-(CH<sub>2</sub>)<sub>p</sub>-

-CH2(CH2)p-

-O-CH=CH-

-NH-CH=CH-90

-N-CH=CH-

wherein p is 2, 3 or 4 and the O and N atoms always 95 are attached to position 4 in the pyridine ring; and physiologically acceptable salts of the compounds I wherein X is S; with the provisos that

(a) not more than one of R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> is hydrogen,

(b) when X is SO, R5 is H and R6, R7 and R8 are 100 selected only from hydrogen, methyl, methoxy, ethoxy, methoxyethoxy and ethoxyethoxy and at the same time more than one of R1, R2, R3 and R4 are hydrogen, then those radicals R1, R2, R3 and R4 which 105 are not H cannot be selected only from alkyl groups,

halogen, alkoxycarbonyl, alkoxy or alkanoyl.

(c) when X is S, R<sup>5</sup> is H, alkanoyl or alkoxycarbonyl, and  $R^6$ ,  $R^7$  and  $R^8$  are selected only from hydrogen, methyl, ethyl, methoxy, ethoxy, methoxyethoxy and 110 ethoxyethoxy and at the same time more than one of  $R^1$ ,  $R^2$ ,  $R^3$  and  $R^4$  are hydrogen, then those radicals  $R^1$ , R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> which are not H cannot be selected only from alkyl groups, halogen, alkoxycarbonyl, alkoxy, alkanoyl, trifluoromethyl, or NO2,

(d) when X is SO, one of  $R^6$ ,  $R^7$  and  $R^8$  is H and the other two of R6, R7 and R8 are alkyl, and at the same time more than one of R1, R2, R3 and R4 are hydrogen, then those radicals R1, R2, R3 and R4 which are not H cannot be sele sted only from alkyl, halogen, cyano,

O O

-C-(alkoxy), (alkyl)-OC-(alkyl)-, alkoxy, hydroxyalkyl,
O

CF<sub>3</sub> or (alkyl)-C-

(e) when R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup> and R<sup>15</sup> are H and simultaneousby R<sup>6</sup> and R<sup>8</sup> are H or CH<sub>3</sub> and R<sup>7</sup> is OCH<sub>3</sub>, then R<sup>1</sup> is not 5 CF<sub>3</sub> when R<sup>2</sup> is H, and R<sup>2</sup> is not CF<sub>3</sub> when R<sup>1</sup> is H.

A compound according to claim 1 wherein X=S.

3. A compound according to claim 1 wherein X=SO.

 4. A compound according to any one of the preceding claims wherein R<sup>5</sup>=H.

5. A compound according to any one of the preceding claims wherein R<sup>15</sup>=H.

 A compound according to any one of the
 preceding claims wherein at least three of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are other than hydrogen, or they form at least one ring.

A compound according to any one of the preceding claims wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are
 selected from H, alkyl and alkoxy groups.

8. A compound according to any one of the preceding claims wherein  $R^{\delta}$  and  $R^{\delta}$  are selected from H, CH<sub>3</sub>, C<sub>2</sub>H<sub>5</sub>, C<sub>3</sub>H<sub>7</sub>, CH(CH<sub>3</sub>)<sub>2</sub> and ring structures connecting with position 4 in the pyridine ring.

 9. A compound according to any one of the preceding claims wherein two of the radicals R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> form one ring structure and the third radical of R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> is H or alkyl.

A compound according to any one of claims
 1-8 wherein R<sup>5</sup> and R<sup>15</sup> are H; at least three of the radicals R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> are other than H; R<sup>6</sup> and R<sup>8</sup> are each H or CH<sub>3</sub>; and R<sup>7</sup> is CH<sub>3</sub>, OCH<sub>3</sub> or OCH<sub>2</sub>CH=CH<sub>2</sub>.

11. A compound of the formula:

35 wherein X is S or SO

R2 is CH3, C2H5, CH(CH3)2 or OCH3.

12. A process for the preparation of a compound of the formula:

wherein  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ ,  $R^6$ ,  $R^7$ ,  $R^8$  and  $R^{15}$  are as 40 defined in claim 1, and X is SO

by oxidizing a compound of the formula l,

wherein R<sup>15</sup>, R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> have the meanings given above, to give a compound of the same formula I wherein X is S0;

13. Process for preparation of a compound of the formula I wherein R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup>, R<sup>4</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup>, R<sup>6</sup> and R<sup>15</sup> are as defined in claim 1 and X is S by reacting a compound of the formula:

$$\begin{array}{c|c}
R^2 & R^1 \\
R^3 & R^4 & R^5
\end{array}$$

50 with a compound of the formula:

$$z^{2} \xrightarrow[l]{R^{5}} R^{5}$$

in which formulae  $R^{15}$ ,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^5$ ,  $R^6$ ,  $R^7$  and  $R^8$  are as defined in claim 1 and wherein one of  $Z^1$  and  $Z^2$  is SH and the other is a leaving group, to give a compound of the formula I wherein X is S.

14. Process for the preparation of a compound of the formula I wherein X is S and at least one of R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> is an ester group (Z)<sub>n</sub>-A-COOR<sup>9</sup>, COOR<sup>10</sup> or (Z)<sub>n</sub>-A-OCOR<sup>10</sup> wherein Z, n, A, R<sup>9</sup> and R<sup>10</sup> are as defined in claim 1 by esterification of a compound of the formula:

wherein R<sup>15</sup>, R<sup>5</sup>, R<sup>6</sup>, R<sup>7</sup> and R<sup>8</sup> are as defined in claim 1 and Y<sup>1</sup>, Y<sup>2</sup>, Y<sup>3</sup> and Y<sup>4</sup> represent either R<sup>1</sup>, R<sup>2</sup>, R<sup>3</sup> and R<sup>4</sup> as defined in claim 1, respectively, or the groups (Z)<sub>n</sub>-A-COOH, COOH and (Z)<sub>n</sub>-A-OH, but at least one of Y<sup>1</sup>, Y<sup>2</sup>, Y<sup>3</sup>, Y<sup>4</sup> is in the acid or alcohol form, by reaction with the appropriate alcohol R<sup>9</sup>OH, R<sup>10</sup>OH or carboxylic acid R<sup>10</sup>COOH, respectively, to form the required compound.

15. Process for preparation of a compound of the formula I wherein R<sup>5</sup> is R<sup>14</sup>CO and R<sup>14</sup> is as defined in claim 1, by acylation of a compound of the formula:

wherein  $R^{15}$ , X,  $R^1$ ,  $R^2$ ,  $R^3$ ,  $R^4$ ,  $R^6$ ,  $R^7$  and  $R^8$  are as defined in claim 1, by reaction with an appropriate acylating agent (R14CO)2O, or R14COX1, wherein X1 is a leaving group.

16. Process for the preparation of a compound of the formula I wherein R5 is H, by hydrolyzing a compound of the formula

$$R^{\frac{1}{2}} \bigvee_{N=1}^{R^{\frac{1}{2}}} \bigcap_{\substack{N=1\\N\\15}}^{R^{\frac{1}{2}}} \bigvee_{N=1}^{R^{\frac{1}{2}}} \bigcap_{N=1}^{R^{\frac{1}{2}}} \bigcap_{N=1}^{R^{\frac{1}{2}}}$$

wherein X, R15, R1, R2, R3, R4, R6, R7 and R8 are as defined in claim 1 and Z3 is a suitable N-protecting 10 group to form the required compound.

- 17. A process according to any one of claims 13-16 wherein a compound in which X is S is obtained and the resulting compound is converted into a physiologically acceptable salt.
- 18. A process according to any one of claims 12-17 substantially as hereinbefore described with reference to any one of the Examples.
- 19. A pharmaceutical composition containing a compound or salt according to any of claims 1-11 20 together with an inert carrier or diluent.
  - 20. A composition according to claim 19 substantially as hereinbefore described with reference to any one of Examples 167-169.
- 21. A compound according to any one of claims 25 1-11 or a physiologically acceptable salt thereof or a composition according to claim 19 or 20 for use in a method of treatment of the human or animal body by surgery or therapy.
- 22. A compound according to any one of claims 30 1-11 or a physiologically acceptable salt thereof or a composition according to claim 19 or 20 for use in the treatment of gastric disorders.
- 23. A compound as defined in any of claims 1-11, or a therapeutically acceptable salt thereof, or a 35 composition according to claim 19 or 20 for use in inhibiting gastric acid secretion in the human or apimal body.
- 24. A compound as defined in any of claims 1-11, or a therapeutically acceptable salt thereof, or a 40 composition according to claim 19 or 20 for use as a gastrointestinal cytoprotecting agent in the human or animal body.
- 25. A compound as defined in any of claims 1-11, or a therapeutically acceptable salt thereof, or a 45 composition according to claim 19 or 20 for use in the treatment of gastrointestinal inflammatory diseases in the human or animal body.
  - 26. A compound of the formula:

$$R^{2a} \xrightarrow{R^{1a}} N z^{1a}$$
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wherein  $R^{1a}$ ,  $R^{2a}$ ,  $R^{3a}$  and  $R^{4a}$  are the same or different 50 and selected from the groups

- (a) H,
- (b) alkyl containing 1-6 carbon atoms including cycloalkyl
- (c) alkoxyalkyl containing 1-3 carbon atoms in the 55 alkoxy residue and 1-6 carbon atoms in the alkyl residue,
  - (d) aryloxyalkyl containing 1-6 carbon atoms in the alkyl residue.
- (e) arylalkyl containing 1-6 carbon atoms in the 60 alkyl residue,
  - (f) aryl,
  - (g) alkoxy containing 1-6 carbon atoms,
- (h) alkoxyalkoxy containing 1-3 carbon atoms in the outer alkoxy residue and 1-6 carbon atoms in the 65 alkoxy residue nearest the aromatic ring.
  - (i) aryloxyalkoxy containing 1-6 carbon atoms in the alkoxy residue,
  - (j) arylalkoxy containing 1-6 carbon atoms in the alkoxy residue, and
  - (k) aryloxy, R<sup>5a</sup> is (a) H,
    - (b) alkoxycarbonyl containing 1-4 carbon atoms in
    - the alkoxy residue, (c) arylalkoxycarbonyl containing 1-2 carbon
- 75 atoms in the alkoxy residue,
  - (d) dialkylaminocarbonyl containing 1-4 carbon atoms in each alkyl residue, or
  - (e) arylaminocarbonyl, and Z1ª is (a) SH,
  - (b) ClorBr provided that not more than one of  $R^{1\text{\tiny o}}, R^{2\text{\tiny o}}, R^{3\text{\tiny o}}$  and R4 is H.
    - 27. A compound of the formula:

wherein R60 and R50 are

- (a) Hor
- (b) alkyl containing 1-5 carbon atoms, and R7ª is (a) alkenyloxy containing 2-5 carbon atoms,
  - (b) alkynyloxy containing 2-5 carbon atoms,
- (c) oxacycloalkyl containing one oxygen atom and 3-7 carbon atoms.
  - (d) oxacycloalkoxy containing two oxygen atoms and 4-7 carbon atoms,
  - (e) oxacycloalkylalkyl containing one oxygen atom
- 95 and 4-7 carbon atoms
  - (f) oxacycloalkylalkoxy containing two oxygen
  - atoms and 4-6 carbon atoms, or
- (g)  $R^{6a}$  and  $R^{7a}$ , or  $R^{7a}$  and  $R^{8a}$  together with the adjacent carbon atoms in the pyridine ring form a ring 100 wherein the part constituted by R<sup>6a</sup> and R<sup>7a</sup> or R<sup>7a</sup> and R80 is
  - -CH=CH--CH=CH--
  - —O—(CH2)00—
  - -CH2-(CH2)pa-
- \_O\_CH=CH\_

wherein pais 2, 3 or 4 and the O atom always is attached to position R70, and Z2 is (a) SH,

(b) halogen CI, Br, 1 or

# (c) OH

provided that not more than one of R<sup>6e</sup> and R<sup>8e</sup> is H.

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